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Finite Element Analysis for Degenerative Cervical Myelopathy: Scoping Review of the Current Findings and Design Approaches, Including Recommendations on the Choice of Material Properties

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Abstract

Background: Degenerative cervical myelopathy (DCM) is a slow-motion spinal cord injury caused via chronic mechanical loading by spinal degenerative changes. A range of different degenerative changes can occur. Finite element analysis (FEA) can predict the distribution of mechanical stress and strain on the spinal cord to help understand the implications of any mechanical loading. One of the critical assumptions for FEA is the behavior of each anatomical element under loading (ie, its material properties).

Objective: This scoping review aims to undertake a structured process to select the most appropriate material properties for use in DCM FEA. In doing so, it also provides an overview of existing modeling approaches in spinal cord disease and clinical insights into DCM.

Methods: We conducted a scoping review using qualitative synthesis. Observational studies that discussed the use of FEA models involving the spinal cord in either health or disease (including DCM) were eligible for inclusion in the review. We followed the PRISMA-ScR (Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews) guidelines. The MEDLINE and Embase databases were searched to September 1, 2021. This was supplemented with citation searching to retrieve the literature used to define material properties. Duplicate title and abstract screening and data extraction were performed. The quality of evidence was appraised using the quality assessment tool we developed, adapted from the Newcastle-Ottawa Scale, and shortlisted with respect to DCM material properties, with a final recommendation provided. A qualitative synthesis of the literature is presented according to the Synthesis Without Meta-Analysis reporting guidelines.

Results: A total of 60 papers were included: 41 (68%) “FEA articles” and 19 (32%) “source articles.” Most FEA articles (33/41, 80%) modeled the gray matter and white matter separately, with models typically based on tabulated data or, less frequently, a hyperelastic Ogden variant or linear elastic function. Of the 19 source articles, 14 (74%) were identified as describing the material properties of the spinal cord, of which 3 (21%) were considered most relevant to DCM. Of the 41 FEA articles, 15 (37%) focused on DCM, of which 9 (60%) focused on ossification of the posterior longitudinal ligament. Our aggregated results of DCM FEA indicate that spinal cord loading is influenced by the pattern of degenerative changes, with decompression alone (eg, laminectomy) sufficient to address this as opposed to decompression combined with other procedures (eg, laminectomy and fusion).

Conclusions: FEA is a promising technique for exploring the pathobiology of DCM and informing clinical care. This review describes a structured approach to help future investigators deploy FEA for DCM. However, there are limitations to these recommendations and wider uncertainties. It is likely that these will need to be overcome to support the clinical translation of FEA to DCM.

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KEYWORDS
scoping review; fine element analysis; cervical spine; spinal cord; degenerative cervical myelopathy

Introduction

Degenerative cervical myelopathy (DCM) occurs when arthritic changes to the structure of the cervical spine injure the spinal cord, causing a slowly progressive spinal cord injury (SCI) [1]. This leads to a range of different symptoms that can affect the whole body, including loss of dexterity, imbalance, altered sensation, bladder and bowel dysfunction, and pain [2]. Although DCM is estimated to affect 1 in 50 adults, <20% are estimated to receive a diagnosis. This is likely, in part, as most are only mildly affected [3,4]. Treatment is currently limited to surgery but, due to inherent risks, is reserved for those with progressive or moderate-to-severe disease [5]. Notably, <5% of patients with DCM will make a complete recovery after surgery, and instead are left with lifelong disabilities and dependence having among the lowest quality of life scores of any disease [6,7]. Consequently, this was recently estimated to cost GBP £0.7 billion (approximately US $0.9 billion) per year [8].

The etiology and pathophysiology of DCM are poorly understood [1,9]. At a macroscopic level, this is a cohort that displays progressive cervical myelopathy with degenerative changes to the structure of their cervical spine, typically causing some deformation of the spinal cord on magnetic resonance imaging (MRI), which responds to decompressive surgery. This led to the hypothesis that DCM is triggered by a chronic mechanical injury, specifically compression loading. However, this is likely to be an oversimplification. Spinal cord compression is most commonly an incidental finding [3]; the amount of compression visualized on the MRI poorly correlates with the disease severity and does not predict the treatment response [10-12]. Moreover, many other forms of mechanical loading also occur, including stretching or shear loading. These are recognized to be capable of causing tissue injury independently [1]. For example, stretching is considered the etiology of myelopathy in tethered cord syndrome and some forms of deformity [13]. Consequently, it is more likely that the mechanical trigger in DCM is the interaction of these mechanical forces rather than one alone. As the structural changes within the spine highly vary between patients, this is likely to be a very individualized phenomenon [14]. This presents a problem for clinical practice, as conventional diagnostic tests such as MRI cannot measure mechanical stress; however, the goal of surgery is to alleviate it [12,15].

Finite element analysis (FEA) is an engineering technique that uses a computational model to derive the extent and severity of mechanical stress from an assumed loading [16]. This has frequently been applied to health care, including, to some extent, SCI and, more recently, DCM [16-18]. FEA could have important applications in DCM, both to improve our understanding of the pathobiology and to represent an individual’s injury and objectively inform surgical strategy. To perform an FEA, a computer model incorporating the geometry, motion, and material properties of each structure must be created [17]. Geometry and motion, to a large extent, can be defined based on an individual’s clinical imaging. However, the material properties must be chosen from other sources. These choices will influence the results of the FEA. For spinal cord FEA to date, these choices have been made on a project-by-project basis, typically informed by the experience of the investigators, their interpretation and knowledge of the literature, and their specific project aims. To inform the development of FEA for DCM, we adopted an iterative approach using a scoping review methodology with the following aims:

- To describe how FEA models have been constructed with respect to spinal cord disease
- To identify and appraise the experimental literature that has informed their material property choices to make recommendations on the material properties for DCM FEA
- To aggregate the findings from studies using FEA to explore DCM.

To the best of our knowledge, this represents a unique approach to selecting the material properties for a clinical FEA model and may represent an exemplar for similar initiatives.

Methods

A scoping review methodology was considered most appropriate to meet these objectives [19]. This scoping review was reported in accordance with the PRISMA-ScR (Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews) guidelines (Multimedia Appendix 1).

Search Strategy

The search was conducted using a modified population, interventions, comparisons, and outcomes strategy, which states that the research question for a review must include the population, intervention, comparison, and outcome. Our research question was, “what are the current findings and design approaches for FEA in DCM?” with the population being patients with DCM, intervention being FEA, and outcomes being current findings and design approaches. To more comprehensively guide future decisions regarding the application of FEA methods to DCM, we broadened our inclusion criteria to incorporate any study that applied FEA to the spinal cord (in either health or disease). Consequently, the search terms were designed to capture observational studies that had developed FEA models that included the spinal cord in either health or disease, including DCM (Multimedia Appendix 2).

Searches were conducted from inception (February 12, 2021) to September 1, 2021, in the MEDLINE and Embase databases. Search sensitivity was evaluated using 5 papers known to meet these objectives [19]. This scoping review was reported in accordance with the PRISMA-ScR (Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews) guidelines (Multimedia Appendix 1).

Inclusion and Exclusion Criteria

Papers were considered eligible for inclusion if they were observational studies that discussed the use of FEA models that
included the spinal cord of humans or animals in either health or disease, including DCM.

Papers were excluded if they were written in a language other than English, did not use FEA models, or did not include the spinal cord in the FEA model. Furthermore, systematic reviews, scoping reviews, editorials, and abstracts were excluded.

**Study Screening and Data Extraction**

Two reviewers (BMD and SS) independently performed title and abstract screening with blinding using Rayyan (Rayyan Systems Inc). A pilot screen of 100 publications was conducted to ensure concordance between reviewers. Any disagreements following unblinding were resolved by discussion between the reviewers until mutual agreement was reached. In this review, papers identified through our search strategy are termed “FEA articles”.

From the included FEA articles, the references used to justify a structure’s material properties were also screened to identify experimental studies reporting original data acquired from physical tissue tests. Studies exploring behavior computationally but including their original physical experiments, even if published elsewhere, were included. Studies that explored properties solely on a computational basis were excluded. This forward search continued within the references of a referenced study if the reference did not meet this criterion and had cited an alternative source.

Papers were retrieved for full-text screening and data extraction using a piloted pro forma. Data extracted from the papers included: author, year of publication, country, study objectives, study design (eg, human or animal study), disease of interest (if any), spinal segment (eg, cervical, thoracic, and lumbar), reference for anatomy (eg, cadaveric specimen and imaging), and details of how the FEA model was developed and validated (including the material properties of the anatomical elements).

Data extraction focused on the properties specifically referenced by the original FEA models and may not have included all the material properties discussed in the paper. To understand an investigator’s approach to model development, these were distinguished as those used to define the model a priori (ie, referenced data and the choice of material law and selected coefficients) or those used to validate the final model (if performed). However, for the purpose of selecting data to inform an FEA model, these references were aggregated and termed as “source articles” in this review.

In the absence of a standard quality assessment tool for experimental studies of biomechanics, we developed a classification to help appraise source articles that are most appropriate for a DCM FEA model [24]. This included a risk of bias assessment adapted from the Newcastle-Ottawa Scale, focusing on selection and reporting bias (Multimedia Appendix 3) [25].

**Data Analysis and Reporting**

Due to significant heterogeneity between methodologies, meta-analysis was not possible, and a qualitative Synthesis Without Meta-Analysis (SWiM) was instead performed. Data were aggregated, where applicable, qualitatively, quantitatively, or using frequency statistics, as per the SWiM guidelines [26].

Given the small field size, with many papers published by single groups, citation networks were created to graphically consider which choices were made across the field and how they were informed. Using this framework and our judgment, we ranked source articles into approximate tertiles. For FEA articles that had cited top-source articles and represented the material properties using an equation, the performance of this equation was further evaluated graphically by generating stress-strain curves. These were exclusively either linear or hyperelastic.

For models using a linear elastic equation, the Young modulus was used as the gradient of the stress-strain curve. For models using a hyperelastic equation, a 3x3 element cube was created using ABAQUS (Dassault Systèmes). The cube was stretched uniaxially, with no constraint applied in the orthogonal directions, linearly increasing the nominal strain in increments of 0.04 to a maximum of 0.4. The outputs of this model were then applied true stress as a function of the applied true strain. Finally, any primary clinical papers that conducted FEA for the investigation of DCM were aggregated separately and analyzed.

Data were displayed using a range of plots constructed using R Studio (version 4.0.3; Posit).

**Results**

**Overall Approach of FEA Models of Spinal Cord Disease: Anatomy, Geometry, Motion, and Validation**

The search returned 597 articles, of which 155 (25.9%) were duplicates (Figure 1). Following screening, 41 FEA articles were eligible for inclusion, of which 32 (78%) modeled the human spinal cord; a further 45 (7.54%) source articles were identified through citation search, of which 19 (42%) were shortlisted as suitable. Of the FEA articles, approximately half (21/41, 51%) focused on SCI [27-47]; 34% (14/41) on DCM [18,20-22,48-57]; and 5% (2/41) each on scoliosis [58,59], syringomyelia [60,61], and flexion myelopathy [62,63]. Most models (25/41, 61%) included only the spinal cord, whereas 24% (10/41) included the surrounding anatomy at multiple vertebral levels, and 17% (7/41) included the surrounding anatomy at only 1 motion segment (ie, 2 adjacent vertebrae). Physiological movement of the spine (flexion and extension) was incorporated into 17% (7/41) of the models, but none evaluated spinal cord oscillation. This was equally likely among the DCM and SCI models (Multimedia Appendix 4).

The anatomy of each model was built using a combination of imaging and cadaveric data in 27% (11/41) of the FEA articles. Typically, imaging was used for bones and cadavers for soft tissues, including the spinal cord. This included an open-source reference library called BodyWorks [64] and a review of spinal cord geometry [65]. MRI was used to define the spinal cord specifically in 20% (8/41) of the FEA articles.
For most FEA articles (33/41, 80%), the spinal cord was modeled as gray matter and white matter separately and had a defined pial layer (26/41, 63%) or was encased within the dural layer (26/41, 63%). Defined pial and dural layers were used in combination in only half of these articles (13/41, 32%). Cerebrospinal fluid (CSF) was specifically modeled in 41% (17/41) of the FEA articles, while other elements were variably included. This choice was independent of the disease and publication date (Multimedia Appendix 4). Elements were modeled using solid shell elements, unless specified differently in the Material Properties of Anatomical Elements With Recommendations for DCM FEA section.

Validation methods were specified in 63% (26/41) of the FEA articles, with 15% (6/41) using their own experiments and 9% (2/41) using literature (Multimedia Appendix 5). These references pointed to 17 articles, of which 7 (42%) provided material property data for the spinal cord in healthy circumstances and 3 (18%) in traumatic SCI circumstances. Of the remaining 17 articles, 4 (24%) described motion of the spine [66-69] and 1 (6%) described the spinal cord in flexion and extension [70]. Of the 9 articles providing information on healthy spinal cord properties, 7 (78%) were also used in other studies to inform the selection of material property. No DCM-specific validation data sets were identified.

**Material Properties of Anatomical Elements With Recommendations for DCM FEA**

**Spinal Cord**

The material properties of the whole spinal cord were defined in 22% (9/41) of the FEA articles. This was rarely justified, but if so, qualified by its uncertain significance [71,72]. Typically, a hyperelastic Ogden variant (4/9, 44%) or a linear elastic (3/9, 33%) function was used.

For the remaining models, gray and white matter were modeled separately, except for the article that explored the impact of a range of white matter material properties, where the material law applied to gray matter was the same as that of white matter. The remaining 32 models were mostly based on tabulated data from the studies by Ichihara et al [72,73], and less frequently, Bilston and Thibault [74], Tunturi [75], and Ozawa et al [76]. Alternatively, a hyperelastic Ogden variant (10/41, 24%) or a linear elastic (4/41, 10%) function was used.

A total of 2 studies specifically compared different material properties with respect to a transverse contusion model of SCI. Jannesar et al [38] explored white matter properties on the basis that single constitutive models may not account for the dynamic (viscoelastic) and anisotropic properties. They identified that this could be improved by adding reinforcing functions. A second order reduced polynomial hyperelastic function combined with a quadratic reinforcing function in a 4-term Prony series performed best ($0.89 < R^2 < 0.99$), although this was principally in relation to the high strain rates of an SCI. Fournely et al [45] used a first-order Ogden function but varied the stiffness of the gray matter with respect to the white matter. Although this fell within the range of the validation data set, they observed differing responses to the load. When the gray matter was stiffer than the white matter, strain distribution was more diffuse and maximal within the white matter. When the stiffness was equivalent, strain was localized to the impact site. When the white matter was stiffer than the gray matter, strain was less localized, maximal within the gray matter and involved the contralateral gray matter. This was the principal factor determining behavior, ahead of other factors explored, including spinal cord diameter, curvature, and impactor angle.

A total of 2 studies similarly explored the implications of different gray and white matter material properties with respect...
to DCM, with similar findings discussed in the Findings From the FEA Studies of DCM section [34,50].

A total of 14 source articles were identified describing the material properties of the spinal cord or its subcomponents (Multimedia Appendix 6 [46,72-75,77-90]), of which 3 (21%) were shortlisted with relevance to an FEA for DCM [72-74]. Their interpretations varied across studies (Figure 2). The choice of material laws and values of those who directly cited the prioritized source articles and separately distinguished gray and white matter are listed in Tables 1-2. Broadly, these align with the source articles; however, there are differences across the strain range (Multimedia Appendix 6). Of these FEA articles representing material properties with an equation, studies by Jannesar et al [29] and Khuyagbaatar et al [53] were selected as these were most aligned for gray matter and white matter, respectively.

**Figure 2.** Network analysis of finite element analysis models, which is linked to a shortlisted source article, for the white matter (A) and gray matter (B) or the spinal cord as a whole (C). The original finite element analysis models are represented by their choice of material law as a star (linear elastic), square (hyperelastic), diamond (tabulated), or triangle (other) and their disease of interest as degenerative cervical myelopathy (DCM; red), spinal cord injury (SCI; blue), or other (green). These link to the primary source articles (dots). An intermediate article, that is, the one that did not include primary experimental data, is pale gray. A shortlisted source article is black. Each figure is additionally available as an interactive file; refer to Multimedia Appendix 7. The higher resolution version of this figure is available in Multimedia Appendix 8.
### Table 1. Extracted material equations for the gray matter.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>E (^a) (MPa)</th>
<th>v (^b)</th>
<th>α (^c)</th>
<th>μ (^d) (MPa)</th>
<th>D (^e) (MPa(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jannesar et al [29], 2021</td>
<td>SCI</td>
<td>Ichihara et al [72], 2003</td>
<td>Hyperelastic</td>
<td>Ogden, first Order</td>
<td>—</td>
<td>0.49</td>
<td>10.57</td>
<td>0.0445</td>
<td>0.905 (^h)</td>
</tr>
<tr>
<td>Khuyagbaatar et al [53], 2017</td>
<td>DCM</td>
<td>Ichihara et al [73], 2001</td>
<td>Hyperelastic</td>
<td>Ogden, first Order</td>
<td>—</td>
<td>0.45</td>
<td>14.7</td>
<td>0.0041</td>
<td>50.5</td>
</tr>
<tr>
<td>Jannesar et al [38], 2016</td>
<td>SCI</td>
<td>Ichihara et al [72], 2003</td>
<td>Hyperelastic</td>
<td>Ogden, first Order</td>
<td>—</td>
<td>0.45</td>
<td>7.52</td>
<td>0.0306</td>
<td>6.77</td>
</tr>
<tr>
<td>Khuyagbaatar et al [39], 2016</td>
<td>SCI</td>
<td>Ichihara et al [73], 2001</td>
<td>Hyperelastic</td>
<td>Ogden, first Order</td>
<td>—</td>
<td>0.45</td>
<td>14.7</td>
<td>0.0041</td>
<td>50.5</td>
</tr>
<tr>
<td>Czyz et al [42], 2008</td>
<td>SCI</td>
<td>Ichihara et al [72], 2003</td>
<td>Linear elastic</td>
<td>—</td>
<td>0.656</td>
<td>0.499</td>
<td>—</td>
<td>0.2188</td>
<td>—</td>
</tr>
<tr>
<td>Maikos et al [43], 2008</td>
<td>SCI</td>
<td>Bilston and Thibault [74], 1996</td>
<td>Hyperelastic</td>
<td>Ogden, first Order</td>
<td>—</td>
<td>0.45</td>
<td>4.7</td>
<td>0.0320</td>
<td>6.47</td>
</tr>
<tr>
<td>Scifert et al [44], 2002</td>
<td>SCI</td>
<td>Bilston and Thibault [74], 1996</td>
<td>Linear elastic</td>
<td>—</td>
<td>0.0667</td>
<td>0.499</td>
<td>—</td>
<td>0.0222</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^a\)E: Young modulus.  
\(^b\)v: Poisson ratio. Where missing, the value of v was assumed to be 0.45.  
\(^c\)α: material exponent parameter.  
\(^d\)μ: ground shear hyperelastic modulus.  
\(^e\)D: compressibility constant.  
\(^f\)The single preferred source of the authors based on modeling (Multimedia Appendix 6), where a range of equations were put forward.  
\(^g\)SCI: spinal cord injury.  
\(^h\)Not available.  
\(^i\)Denotes a suspected error in original text and input value given.  
\(^j\)Values in italics are input based on the identity for isotropic materials, D=3(1-2v)/μ(1+ν), and for linear elastic, μ=E/(2(1+v)).  
\(^k\)DCM: degenerative cervical myelopathy.
Table 2. Extracted material equations for the white matter.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>E&lt;sup&gt;a&lt;/sup&gt; (MPa)</th>
<th>ν&lt;sup&gt;b&lt;/sup&gt;</th>
<th>α&lt;sup&gt;c&lt;/sup&gt;</th>
<th>μ&lt;sup&gt;d&lt;/sup&gt; (MPa)</th>
<th>D&lt;sup&gt;e&lt;/sup&gt; (MPa&lt;sup&gt;-1&lt;/sup&gt;)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liang et al [48], 2021</td>
<td>DCM&lt;sup&gt;f&lt;/sup&gt;</td>
<td>Ichihara et al [73], 2001</td>
<td>Linear elastic</td>
<td>➥&lt;sup&gt;g&lt;/sup&gt;</td>
<td>4.2</td>
<td>0.45</td>
<td>—</td>
<td>1.448&lt;sup&gt;b&lt;/sup&gt;</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [52], 2017</td>
<td>DCM</td>
<td>Ichihara et al [73], 2001</td>
<td>Hyperelastic</td>
<td>Ogden, first order</td>
<td>—</td>
<td>0.45</td>
<td>12.5</td>
<td>0.0040</td>
<td>51.7</td>
</tr>
<tr>
<td>Khuyagbaatar et al [39], 2016</td>
<td>SCI&lt;sup&gt;i&lt;/sup&gt;</td>
<td>Ichihara et al [73], 2001</td>
<td>Hyperelastic</td>
<td>Ogden, first order</td>
<td>—</td>
<td>0.45</td>
<td>12.5</td>
<td>0.0040</td>
<td>51.7</td>
</tr>
<tr>
<td>Czyz et al [42], 2008</td>
<td>SCI</td>
<td>Ichihara et al [72], 2003</td>
<td>Linear elastic</td>
<td>—</td>
<td>0.277</td>
<td>0.499</td>
<td>—</td>
<td>0.0924</td>
<td>—</td>
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<tr>
<td>Maikos et al [43], 2008</td>
<td>SCI</td>
<td>Bilston and Thibault [74], 1996</td>
<td>Hyperelastic</td>
<td>Ogden, first order</td>
<td>—</td>
<td>0.45</td>
<td>4.7</td>
<td>0.0320</td>
<td>6.47</td>
</tr>
<tr>
<td>Scifert et al [44], 2002</td>
<td>SCI</td>
<td>Bilston and Thibault [74], 1996</td>
<td>Linear elastic</td>
<td>—</td>
<td>0.0667</td>
<td>0.499</td>
<td>—</td>
<td>0.0222</td>
<td>—</td>
</tr>
</tbody>
</table>

<sup>a</sup>E: Young modulus.<br>
<sup>b</sup>v: Poisson ratio. Where missing, the value of ν was assumed to be 0.45.<br>
<sup>c</sup>α: material exponent parameter.<br>
<sup>d</sup>μ: ground shear hyperelastic modulus.<br>
<sup>e</sup>D: compressibility constant.<br>
<sup>f</sup>DCM: degenerative cervical myelopathy.<br>
<sup>g</sup>Not available.<br>
<sup>i</sup>SCI: spinal cord injury.

Pia

Of the 26 FEA articles with defined pia, 14 (54%) used a linear elastic function, 9 (21%) did not report their method, and 2 (5%) used a hyperelastic Ogden variant function. The remaining study (1/26, 4%) used tabulated data from the study by Ichihara et al [73].

A total of 4 source articles were identified for the pia (Multimedia Appendix 6), of which 2 (50%) were shortlisted as suitable [75,77]. The choice of material laws and the values of those who directly cited these shortlisted source articles are listed in Table 3. These equations have differences in how they represent the source article (Multimedia Appendix 6). Of the FEA articles representing material properties with an equation, the study by Jannesar et al [38] was selected as the most preferred.
### Table 3. Extracted material equations for the pia.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>( E^a ) (MPa)</th>
<th>( \nu^b )</th>
<th>( \alpha^c )</th>
<th>( \mu^d ) (MPa)</th>
<th>( D^e ) (MPa(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Henao et al [58,59], 2017</td>
<td>Other DCM</td>
<td>Tunturi [75], 1978</td>
<td>Linear elastic</td>
<td>—</td>
<td>—</td>
<td>100</td>
<td>0.4</td>
<td>—</td>
<td>35.71</td>
</tr>
<tr>
<td>Nishida et al [91], 2016</td>
<td>DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Nishida et al [54], 2015</td>
<td>DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Nishida et al [55], 2014</td>
<td>DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Nishida et al [22], 2012</td>
<td>DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Henao et al [58], 2018</td>
<td>Other DCM</td>
<td>Tunturi [75], 1978</td>
<td>Linear elastic</td>
<td>—</td>
<td>—</td>
<td>100</td>
<td>0.4</td>
<td>—</td>
<td>35.71</td>
</tr>
<tr>
<td>Kato et al [56], 2010</td>
<td>DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Kato et al [62], 2008</td>
<td>Other DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Kato et al [63], 2009</td>
<td>Other DCM</td>
<td>Tunturi [75], 1978</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Jannesar et al [38], 2016</td>
<td>SCI</td>
<td>Kimpara et al [77], 2006</td>
<td>Linear elastic</td>
<td>—</td>
<td>—</td>
<td>39.3</td>
<td>0.3</td>
<td>—</td>
<td>15.12</td>
</tr>
</tbody>
</table>

\( a \): Young modulus.

\( b \): Poisson ratio. Where missing, the value of \( \nu \) was assumed to be 0.45.

\( c \): Material exponent parameter.

\( d \): Ground shear hyperelastic modulus.

\( e \): Compressibility constant.

\( f \): Not available.

\( g \): Values in italics are input based on the identity for isotropic materials, \( D = 3(1-2\nu)/(\mu(1+\nu)) \), and for linear elastic, \( \mu = E/(2(1+\nu)) \).

\( h \): DCM: degenerative cervical myelopathy.

\( i \): The single preferred source of the authors based on modelling (Multimedia Appendix 6).

### Dura

Of the 26 models with defined dura, 18 (69%) used a linear elastic function, 5 (19%) used a hyperelastic Ogden variant, and 3 (12%) did not report their method.

Persson et al [46] compared the performance of a linear and hyperelastic function, which is summarized in the following CSF section.

A total of 9 source articles were referenced (Multimedia Appendix 6), of which 4 (44%) were shortlisted [78-81]. The choice of material laws and values of those who directly cited these prioritized source articles are listed in Table 4. These equations have differences in how they represent the source article (Multimedia Appendix 6). Of the FEA articles representing material properties with an equation, the study by Sparrey et al [33] was selected as preferred.
Table 4. Extracted material equations for the dura.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>$E^a$ (MPa)</th>
<th>$\nu^b$</th>
<th>$\alpha^c$</th>
<th>$\mu^d$ (MPa)</th>
<th>$D^e$ (MPa$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stoner et al [20], 2020</td>
<td>DCM$^f$</td>
<td>Persson et al [92], 2020</td>
<td>Linear Elastic $^g$</td>
<td>—</td>
<td>5</td>
<td>0.45</td>
<td>—</td>
<td>1.72 $^h$</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [49], 2018</td>
<td>DCM</td>
<td>Persson et al [92], 2020</td>
<td>Linear Elastic</td>
<td>—</td>
<td>80</td>
<td>0.49</td>
<td>—</td>
<td>26.85</td>
<td>—</td>
</tr>
<tr>
<td>Henao et al [58,59], 2017</td>
<td>Other</td>
<td>Wilcox et al [47], 2004</td>
<td>Linear Elastic</td>
<td>—</td>
<td>231</td>
<td>0.45</td>
<td>—</td>
<td>79.66</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [52], 2017</td>
<td>DCM</td>
<td>Persson et al [92], 2020</td>
<td>Linear Elastic</td>
<td>—</td>
<td>80</td>
<td>0.49</td>
<td>—</td>
<td>26.85</td>
<td>—</td>
</tr>
<tr>
<td>Sparrey et al [33], 2016$^i$</td>
<td>SCI</td>
<td>Hong et al [78], 2011 and Zarzur et al [79], 1996</td>
<td>Hyper-elastic Ogden, 1st Order</td>
<td>—</td>
<td>0.45</td>
<td>16.2</td>
<td>1.205</td>
<td>0.172</td>
<td>—</td>
</tr>
<tr>
<td>Yan et al [36], 2012</td>
<td>SCI</td>
<td>Wilcox et al [47], 2004</td>
<td>Linear Elastic</td>
<td>—</td>
<td>142</td>
<td>0.45</td>
<td>—</td>
<td>48.97</td>
<td>—</td>
</tr>
<tr>
<td>Henao et al [58], 2018</td>
<td>Other</td>
<td>Wilcox et al [47], 2004</td>
<td>Linear Elastic</td>
<td>—</td>
<td>231</td>
<td>0.45</td>
<td>—</td>
<td>79.66</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [39], 2016</td>
<td>SCI</td>
<td>Persson et al [92], 2020</td>
<td>Linear Elastic</td>
<td>—</td>
<td>80</td>
<td>0.49</td>
<td>—</td>
<td>26.85</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [57], 2015</td>
<td>DCM</td>
<td>Persson et al [92], 2020</td>
<td>Linear Elastic</td>
<td>—</td>
<td>80</td>
<td>0.49</td>
<td>—</td>
<td>26.85</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [57], 2015</td>
<td>SCI</td>
<td>Persson et al [92], 2020</td>
<td>Linear Elastic</td>
<td>—</td>
<td>80</td>
<td>0.49</td>
<td>—</td>
<td>26.85</td>
<td>—</td>
</tr>
<tr>
<td>Czyz et al [42], 2008</td>
<td>SCI</td>
<td>Wilcox et al [47], 2004</td>
<td>Linear Elastic</td>
<td>—</td>
<td>142</td>
<td>0.45</td>
<td>—</td>
<td>48.97</td>
<td>—</td>
</tr>
<tr>
<td>Persson et al [46], 2011</td>
<td>SCI</td>
<td>Wilcox et al [47], 2004</td>
<td>Linear Elastic</td>
<td>—</td>
<td>80</td>
<td>0.49</td>
<td>—</td>
<td>26.85</td>
<td>—</td>
</tr>
<tr>
<td>Wilcox et al [47], 2004</td>
<td>SCI</td>
<td>Wilcox et al [47], 2004</td>
<td>Anisotropic Elastic</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

$^a$E: Young modulus.

$^b$ν: Poisson ratio. Where missing, the value of $\nu$ was assumed to be 0.45.

$^c$α: material exponent parameter.

$^d$μ: ground shear hyperelastic modulus.

$^e$D: compressibility constant.

$^f$DCM: degenerative cervical myelopathy.

$^g$Not available.

$^h$Values in italics are input based on the identity for isotropic materials, $D=3(1-2\nu)/(\mu(1+\nu))$, and for linear elastic, $\mu=E/(2(1+\nu))$.

$^i$The single preferred source of the authors based on modelling (Multimedia Appendix 6).

Dentate Ligament

Of the 13 FEA articles that included the dentate ligament, 12 (92%) used a linear elastic function and 1 (8%) used tabulated data. Typically, these were modeled using shell elements (6/13, 46%) with geometric properties, but 8% (1/13) used link elements and 15% (2/13) used spring elements.

A total of 2 source articles were referenced (Multimedia Appendix 6), of which both were shortlisted [75,82]. The choice of material laws and values of those who directly cited these prioritized source articles are listed in Table 5.
Table 5. Extracted material equations for the dentate.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>E(^a) (MPa)</th>
<th>(v^b)</th>
<th>(\alpha^c)</th>
<th>(\mu^d) (MPa)</th>
<th>D(^e) (MPa(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Henao et al [58,59], 2017</td>
<td>Other</td>
<td>Tunturi [75], 1978</td>
<td>Linear elastic</td>
<td>—f</td>
<td>100</td>
<td>0.4</td>
<td>—</td>
<td>35.7</td>
<td>—</td>
</tr>
<tr>
<td>Henao et al [58], 2018</td>
<td>Other</td>
<td>Tunturi [75], 1978</td>
<td>Linear elastic</td>
<td>—</td>
<td>100</td>
<td>0.4</td>
<td>—</td>
<td>35.7</td>
<td>—</td>
</tr>
<tr>
<td>Greaves et al [41], 2008</td>
<td>SCI(^h)</td>
<td>Tunturi [75], 1978</td>
<td>Linear elastic</td>
<td>—</td>
<td>5.8</td>
<td>—</td>
<td>—</td>
<td>2.0</td>
<td>—</td>
</tr>
<tr>
<td>Czyz et al [42], 2008</td>
<td>SCI</td>
<td>Tunturi [75], 1978</td>
<td>Linear elastic</td>
<td>—</td>
<td>100</td>
<td>0.3</td>
<td>—</td>
<td>38.5</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^a\)E: Young modulus.
\(^b\)\(v\): Poisson ratio. Where missing, the value of \(v\) was assumed to be 0.45.
\(^c\)\(\alpha\): material exponent parameter.
\(^d\)\(\mu\): ground shear hyperelastic modulus.
\(^e\)D: compressibility constant.
\(^f\)Not available.
\(^g\)Values in italics are input based on the identity for isotropic materials, \(D=3(1-2v)/(\mu(1+v))\), and for linear elastic, \(\mu=E/(2(1+v))\).
\(^h\)SCI: spinal cord injury.

Cerebrospinal Fluid

Of the 17 models that included CSF, 8 (47%) modeled it as a Newtonian fluid. Alternatives included modeling CSF as a pressurized fluid cavity (1/17, 6%), modeling it as a polynomial equation of state (1/17, 6%), modeling it as smoothed particular hydrodynamics (1/17, 6%), using a hyperelastic Mooney-Rivlin model (3/17, 18%), or using a linear elastic equation (1/17, 6%).

Persson et al [46] and Jones et al [93] specifically explored the implications of including a CSF cavity, with or without the dura. To measure cord deformation, Persson et al [46] used an FEA model with reference to a transverse bovine impaction model of SCI, whereas Jones et al [93] performed their own bovine and surrogate cord experiments. They observed that the presence of CSF reduced stress and strain (Persson et al [46]) on the spinal cord and deformation (Jones et al [93]) in the spinal cord. Persson et al [46] demonstrated this was through a greater longitudinal distribution, particularly when the dura was included and modeled using a hyperelastic Ogden (as opposed to linear elastic) function. Furthermore, Persson et al [46] observed that cord deformation occurred upon contact with the dura (before the CSF between the spinal cord and the dura was redistributed). Jones et al [93] observed that the inclusion of the dura only changed behavior if CSF was also included.

Furthermore, Arhipstov and Marom [31] explored CSF pressure, alongside the presence or absence of epidural fat, using a computational contusion model of SCI based on a thoracic burst fracture. Both CSF and epidural fat were modeled using smoothed particular hydrodynamics. In a model without epidural fat, spinal cord stress and strain increased with increasing CSF pressure. However, in the model with epidural fat, spinal cord stress and strain decreased with increasing CSF pressure.

A total of 5 source articles were referenced (Multimedia Appendix 6), of which 3 (60%) were shortlisted [46,83,84]. The choice of material laws and values of those who directly cited these prioritized source articles are listed in Table 6.

Table 6. Extracted material equations for the cerebrospinal fluid.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Viscosity (Pa/s)</th>
<th>Density (kg/m(^3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Khuyagbaatar et al [52], 2017</td>
<td>DCM(^a)</td>
<td>Bloomfield et al [83], 1998</td>
<td>Newtonian Fluid</td>
<td>0.001</td>
<td>— (^b)</td>
</tr>
<tr>
<td>Arhipstov [31], 2021</td>
<td>SCI(^c)</td>
<td>Persson et al [46], 2011</td>
<td>Polynomial Equation of State</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [39], 2016</td>
<td>DCM</td>
<td>Bloomfield et al [83], 1998, Brydon et al [84], 1995</td>
<td>Newtonian Fluid</td>
<td>0.001</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [39], 2016</td>
<td>SCI</td>
<td>Bloomfield et al [83], 1998, Brydon et al [84], 1995</td>
<td>Newtonian Fluid</td>
<td>0.001</td>
<td>1000</td>
</tr>
<tr>
<td>Khuyagbaatar et al [57], 2015</td>
<td>DCM</td>
<td>Bloomfield et al [83], 1998, Brydon et al [84], 1995</td>
<td>Newtonian Fluid</td>
<td>0.001</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [57], 2015</td>
<td>SCI</td>
<td>Bloomfield et al [83], 1998, Brydon et al [84], 1995</td>
<td>Newtonian Fluid</td>
<td>0.001</td>
<td>—</td>
</tr>
<tr>
<td>Persson et al [46], 2011</td>
<td>SCI</td>
<td>Bloomfield et al [83], 1998</td>
<td>Newtonian Fluid</td>
<td>0.001</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^a\)DCM: degenerative cervical myelopathy.
\(^b\)Not available.
\(^c\)SCI: spinal cord injury.
Posterior Longitudinal Ligament and Ligamentum Flavum

The analysis focused on the posterior longitudinal ligament and ligamentum flavum, given their specific involvement in the pathobiology of DCM. In all 6 instances included, they were included together and modeled in the same manner: using piecewise linear plasticity (2/6, 33%), linear elastic function (2/6, 33%), hyperelastic Ogden variant (1/6, 17%), or tabulated data (1/6, 17%).

A total of 6 source articles were referenced (Multimedia Appendix 6), of which 3 (50%) were shortlisted [85-87]. The choice of material laws and values of those who directly cited these prioritized source articles are listed in Tables 7 and 8.

Table 7. Extracted material equations for the ligamentum flavum.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>$E^a$ (MPa)</th>
<th>$\nu^b$</th>
<th>$\alpha^c$</th>
<th>$\mu^d$ (MPa)</th>
<th>$D^e$ (MPa$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Greaves et al [41], 2008</td>
<td>SCI$^f$</td>
<td>Yoganandan et al 1989 and 2000 [86,87]</td>
<td>Linear elastic $^g$</td>
<td>3.8</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1.3 $^h$</td>
<td>—</td>
</tr>
</tbody>
</table>

$^a$E: Young modulus.
$^b$v: Poisson ratio. Where missing, the value of v was assumed to be 0.45.
$^c$α: material exponent parameter.
$^d$μ: ground shear hyperelastic modulus.
$^e$D: compressibility constant.
$^f$SCI: spinal cord injury.
$^g$Not available.
$^h$Values in italics are input based on the identity for isotropic materials, $D=3(1-2\nu)/(\mu(1+\nu))$, and for linear elastic, $\mu=E/(2(1+\nu))$.

Table 8. Extracted material equations for the posterior longitudinal ligament.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>$E^a$ (MPa)</th>
<th>$\nu^b$</th>
<th>$\alpha^c$</th>
<th>$\mu^d$ (MPa)</th>
<th>$D^e$ (MPa$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Greaves et al [41], 2008</td>
<td>SCI$^f$</td>
<td>Przybylski et al [85], 1996 and Yoganandan 1989 and 2000 [86,87]</td>
<td>Linear elastic $^g$</td>
<td>35.7</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>12.3 $^h$</td>
<td>—</td>
</tr>
</tbody>
</table>

$^a$E: Young modulus.
$^b$v: Poisson ratio. Where missing, the value of v was assumed to be 0.45.
$^c$α: material exponent parameter.
$^d$μ: ground shear hyperelastic modulus.
$^e$D: compressibility constant.
$^f$SCI: spinal cord injury.
$^g$Not available.
$^h$Values in italics are input based on the identity for isotropic materials, $D=3(1-2\nu)/(\mu(1+\nu))$, and for linear elastic, $\mu=E/(2(1+\nu))$.

Spinal Roots

A total of 7 models included spinal nerve roots, of which 2 (29%) distinguished between the intradural and extradural components. These 2 models specifically explored the nature of C5 palsy in relation to surgery for DCM [49,57]. Nerve roots were all modeled with spring elements, either as a spring (5/7, 71%) or with a linear elastic equation (2/7, 29%).

A total of 2 source articles of equivalent quality were referenced (Multimedia Appendix 6) [88,89]. The choice of material laws and values of those who directly cited these prioritized source articles are listed in Table 9.
Table 9. Extracted material equations for the nerve roots.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Ea (MPa)</th>
<th>vb</th>
<th>Spring constant</th>
<th>Mass (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lévy et al [18], 2021</td>
<td>DCMc</td>
<td>Kulkarni [88], 2007</td>
<td>Spring</td>
<td>—d</td>
<td>—</td>
<td>0.133</td>
<td>0.1</td>
</tr>
<tr>
<td>Khuyagbaatar et al [49], 2018</td>
<td>DCM</td>
<td>Singh [89], 2005</td>
<td>Linear Elastic</td>
<td>1.3</td>
<td>0.3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Henao et al [58,59], 2017</td>
<td>Other</td>
<td>Kulkarni [88], 2007</td>
<td>Spring</td>
<td>—</td>
<td>—</td>
<td>0.133</td>
<td>—</td>
</tr>
<tr>
<td>Khuyagbaatar et al [52], 2017</td>
<td>DCM</td>
<td>Singh [89], 2005</td>
<td>Linear Elastic</td>
<td>1.3</td>
<td>0.3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Henao et al [58], 2018</td>
<td>Other</td>
<td>Kulkarni [88], 2007</td>
<td>Spring</td>
<td>—</td>
<td>—</td>
<td>0.133</td>
<td>—</td>
</tr>
</tbody>
</table>

a: Young modulus.

b: Poisson ratio; where missing, vb was assumed to be 0.45. For Kulkarni et al [88], the unit is uncertain, with a range of different units referenced across its citations.
c: DCM: degenerative cervical myelopathy.
d: Not available.

Other Elements

Other elements included in some models were bone (14/41, 34%); intervertebral disks (IVDs; 13/41, 31%); and the remaining spinal ligaments, such as the anterior longitudinal or interspinous ligament.

The bone was generally modeled as a rigid body (8/14, 57%). Of the 8 models, 3 (21%) subdivided the vertebrae into anatomical subcomponents (eg, body, laminae, and spinous process), and 5 (36%) distinguished between cortical and cancellous bone, of which 3 (60%) applied an equation just to the cortical bone (linear elastic in all cases) and 2 (40%) applied a Johnson-Cook or plastic kinematic equation. We found no eligible source articles using our search process.

The IVD were modeled as a single entity in 54% (7/13) of the papers, typically as a rigid body (5/7, 71%) or using a linear elastic equation (2/7, 29%). Alternatively, they were modeled separately as nucleus pulposus and annulus fibrosus. Techniques for the nucleus pulposus included a Mooney-Rivlin model (3/6, 50%), Ogden second-order variant (1/6, 17%), and fluid elements (2/6, 33%). The annulus fibrosus included a Mooney-Rivlin model (2/6, 33%), Ogden second-order variant (1/6, 17%), Ogden third-order variant (1/6, 17%), and linear elastic equation (2/6, 33%).

A total of 3 source articles were found for IVD, and 1 was shortlisted (Multimedia Appendix 6) [90]. The choice of material laws and values of those who directly cited these prioritized source articles are listed in Table 10.

Table 10. Extracted material equations for the intervertebral disc.

<table>
<thead>
<tr>
<th>Study, year</th>
<th>Pathology</th>
<th>Reference</th>
<th>Law</th>
<th>Variant</th>
<th>Ea (MPa)</th>
<th>vb</th>
<th>αc</th>
<th>μd (MPa)</th>
<th>Df (MPa⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Greaves et al [41], 2008</td>
<td>SCIf</td>
<td>Spilker et al [90], 1986</td>
<td>Linear elastic</td>
<td>g</td>
<td>3.4</td>
<td>—</td>
<td>—</td>
<td>1.2b</td>
<td>—</td>
</tr>
</tbody>
</table>

a: Young modulus.

b: Poisson ratio. Where missing, the value of vb was assumed to be 0.45.
c: Material exponent parameter.
d: Ground shear hyperelastic modulus.
e: Compressibility constant.
f: SCI: spinal cord injury.
g: Not available.

Values in italics are input based on the identity for isotropic materials, D=3(1-2ν)/(μ(1+ν)), and for linear elastic, μ=E/(2(1+ν)).

Findings From the FEA Studies of DCM

Of the DCM models, 60% (9/15) specifically focused on ossification of the posterior longitudinal ligament (OPLL), a specific subtype of DCM.

Stress and Static Cord Compression

A total of 8 models explored the relationship between the amount of static spinal cord compression and spinal cord stress. Kato et al [56] and Kim et al [21] used parametric models of the spinal cord to explore the implications of OPLL (anterior) compression at 2 adjacent vertebrae. The model was constrained posteriorly, reflecting the lamina. They found that the stress increased with increasing cord compression, with an apparent exponential relationship. Minimal stress was detected at <40% but dramatically increased at ≥50%. This relationship was replicated by Nishida et al [91] using posterior compression, by Liang et al [48] simulating a disk prolapse, and in a multisegmental model of OPLL by Khuyagbaatar et al [52,57]. Furthermore, it was replicated in cervical spondylosis by Levy et al [18] (Figure 3 [18,21,52,57]).

Maximal stress was observed in the gray matter and, to a lesser extent, in the lateral and posterior funiculus. Nishida et al [91] observed differences in the stress distribution at low compression rates depending on the spinal cord level related to differing morphology; however, beyond a compression rate of 30%, this was consistent (Figure 4).
Okazaki et al [50] explored the implications of spinal cord aging using a parametric model of the spinal cord. The model was given white and gray matter properties based on a young or aged bovine spinal cord specimen. They observed that stress increased under a low amount of anterior compression in the aged spinal cord and was more widely distributed throughout the gray matter and white matter. In contrast, the gray matter was unaffected in the young specimen.

**Figure 3.** Spinal cord compression and spinal cord stress in degenerative cervical myelopathy models. For models tabulating the von Mises stress at different measures of static compression or canal stenosis (n=4) [18,21,52,57], the values were plotted on a line graph with a line of best fit representing the average value (blue).

**Figure 4.** Spinal cord compression and location of spinal cord stress in degenerative cervical myelopathy models. The spinal cord was partitioned, per hemicord, as gray matter and anterior, anterolateral, posterolateral, and posterior white matter. For each study, reporting the cross-sectional distribution of Von Mises stress (n=12) and the location of stress that fell within the top 30% of measured stress was noted. These frequencies were aggregated by compression pattern and displayed for (A) anterior diffuse and static, (B) anterior focal and static, and (C) circumferential and dynamic distribution and location of stress as relative proportions.
**Stress and Dynamic Cord Compression**

Nishida et al [22] used a parametric model to explore the implications of ligamentum flavum buckling in neck extension in the context of cervical stenosis. For this, the spinal cord was restricted posteriorly by the ligamentum flavum and then anteriorly, either by a central curvature (representing a disk prolapse) or a flat lateral or flat cross-sectional constraint (representing the ligament). The amount of ligamentum flavum buckling was measured using a kinematic MRI. Spinal cord stress was observed in all scenarios and was maximal using the flat cross-sectional constraint.

Later, Nishida et al [54] used a parametric model of OPLL to demonstrate that while dynamic and static compression alone could stress the spinal cord, they could also act together, although it was unclear whether this was additive or multiplicative. In dynamic compression alone, stress was more restricted to gray matter.

**Stress and Shape of Cord Compression**

Khuyagbaatar et al [57] and Kim et al [21] did not identify any difference in OPLL shape or type with respect to observed spinal cord stress. Furthermore, in the study by Nishida et al [22], the distribution of stress was broadly comparable across the three scenarios affecting the gray matter and anterior and posterolateral aspects of the white matter tracts. In unilateral compression only, the ipsilateral gray matter was affected. Levy et al [18] explored gradually increasing anterior diffuse (broad-based disk), anterior lateral, and circumferential compression using a static multilevel model. Different phenotypes of stress were observed, including peak stress, point of onset, and rate of increase. The highest stress was observed with an anterior diffuse or circumferential compression (Figure 4).

**Stress and Surgical Decompression**

Khuyagbaatar et al [39] used a multisegmental model to explore the implications of hemilaminectomy, laminectomy, and laminoplasty on spinal cord stress following a 1-, 2-, 3-, or 4-level posterior decompression for continuous OPLL. Stress remained elevated following hemilaminectomy but was low and equivalent between laminectomy and laminoplasty. The postoperative deformity was not modeled.

Nishida et al [55] used a parametric model to explore the implications of alignment following posterior decompression for OPLL. They demonstrated that although stress decreased significantly following decompression, it slightly increased in the anterior funiculus, increasing in the gray matter and posterolateral funiculi with progressive deformity. They subsequently replicated this in a separate analysis [51], demonstrating that kyphosis and increased mobility after decompression would elevate the observed stress.

Khuyagbaatar et al [49,52] explored the effects of laminectomy and laminoplasty, respectively, for the treatment of OPLL using a multisegmental static compression model. They demonstrated that all procedures reduced spinal cord stress significantly (>90%), whether in lordotic (K Line positive) or kyphotic deformity (K Line negative) [94]. However, stress was elevated within the exiting C5 nerve root following laminectomy if there was a kyphotic deformity and lateral-type OPLL following laminoplasty. In both instances, the amount of nerve root stress was related to the amount of anterior compression.

Stoner et al [20] used a multisegmental dynamic model (C2-T1) to explore the implications of multilevel C4-7 cervical spondylosis (anterior disk prolapses and osteophyte formation) treated with C4-7 anterior cervical disectomy and fusion (ACDF), laminoplasty, or ACDF with laminectomy. Notably, all procedures caused stress to increase at adjacent levels above those of healthy controls. However, a stand-alone ACDF caused increased stress within the spinal cord at C3 to a level above that of the preoperative DCM model in flexion.

Where possible, these were aggregated, demonstrating that the spinal cord tolerated significant compression before stress increased exponentially (Figure 3 [18,21,52,57]). Aggregating the distributions of stress observed across studies, based on the nature of compression, demonstrated differing stress distributions (Figure 4). For static and diffuse anterior compression, the bilateral posterior white matter and gray matter were the most affected. For static and focal compression, the anterior white matter and, to a lesser extent, the gray matter were most affected. This was observed bilaterally despite a focal or lateral element. For circumferential compression in a dynamic model, the bilateral gray matter and posterior white matter were the most affected.

**Stress and Tissue Injury**

Notably, although differential patterns of stress were observed throughout these DCM models, the levels remained relatively low (<0.5 MPa). DCM FEA models did not explore the relationship between the observed stress and tissue injury.

**Discussion**

**Overview**

FEA is a promising technique used in DCM, although there remain uncertainties regarding the ideal approach and its clinical interpretation. This review highlights the numerous decisions investigators must make when performing FEA, which can affect findings and underpin the need for a systematic approach, as applied in this study. On the basis of current evidence, we have shortlisted our preferred material property choices for a DCM model and conclude that a distinction between gray and white matter is preferable.

**Principal Findings and Comparison to Prior Work**

A total of 15 studies were identified applying FEA to investigate DCM. The insights from these studies broadly align with the current evidence base. First, the spinal cord can tolerate some compression. This is in keeping with clinical practice, where asymptomatic spinal cord compression is far more common [3], and the amount of cord compression is a poor surrogate for disease severity or progression [1]. Second, the movement of the subaxial cervical spine can augment the stress on the spinal cord. This is in keeping with clinical practice, including the concept of dynamic injury and the proposed role of flexion/extension MRI or electrophysiology [95-98]. Finally,
it demonstrated the significant effectiveness of decompression surgery, regardless of the technique, and the comparatively minor gains of using one technique over the other. This is in keeping with clinical practice, where high-quality comparisons of anterior versus posterior surgery are equivalent, and currently, there is no strong evidence that routine stabilization (eg, instrumented fusion vs laminoplasty vs laminectomy or ACDF vs ACDF with a plate) is required [99-102], all pointing toward the need for a personalized surgical approach [15].

Furthermore, although more nuanced findings were proposed by the identified FEA studies and this would require in vivo corroboration, the application of FEA in DCM appears well founded overall. More widely, it also seems potentially valuable and timely. The pathobiology of DCM is poorly understood, with its investigation being among the top 10 global research priorities [1]. Current preclinical models have many limitations. For example, common recent models use an expandable polymer inserted behind the spinal cord and within the canal to cause cervical myelopathy. Therefore, this does not model anterior compression, nor does it truly represent a chronic injury mechanism. Furthermore, in clinical practice, clinical decisions are based on imperfect tools [103]. For example, structural MRI in a supine position defines the nature of degenerative changes but not if, where, or how an SCI occurs. FEA could change this, particularly given the parallel advances in the automatic segmentation of MRI [12].

Furthermore, while this review highlights that FEA is a versatile technique, investigators must make many decisions regarding how it is applied. These decisions can alter the findings and, therefore, must be carefully considered. At this stage, there seem to be only a few pervasive insights. First, it seems prudent to model the white matter and gray matter separately. Ichihara et al [73] demonstrated that these structures have differing material properties, and how they are defined alters the observed stress and strain. Furthermore, these structures age differently, as shown by Ozawa et al [76]. Histological studies of DCM have shown differing disease features among the white matter and gray matter, with the gray matter being the focus of more significant cellular changes [9]. Moreover, aging is an important factor in DCM, associated with greater disease severity, a greater rate of progression, and poorer response to treatment [104]. There are also early indicators that accelerating aging is a pathological process [1]. Therefore, the observation that the gray matter was unaffected in the younger spinal cord specimen is noteworthy [34,45,50].

Second, while some models have chosen to use linear elastic equations, time-independent hyperelastic models more closely reflected the known material properties of the spinal cord. These, or simply tabulated data, were generally adopted by DCM studies and supported by a single study that evaluated different approaches [38]. Conceptually, taking a more faithful approach to modeling the spinal cord material properties is likely to be more applicable to DCM and its etiology, as contrasted with traumatic SCI, spinal cord stress may be below the limits for tissue injury (eg, asymptomatic spinal cord compression), and above (eg, DCM). It is worth noting that none of these approaches considers the impact of repetitive injuries, and it is likely that time dependence in modeling is relevant [1]. Given the timeline of DCM pathogenesis (years), this is likely beyond the normal material scales.

Finally, similar to DCM, as the stresses involved are well below the elastic limit of the bone, the vertebras can be modeled simply as rigid bodies. The critical aspect for bones is instead the way that their geometry and movement affect the loading on the soft tissues.

However, there remain many uncertainties for further evaluation. These include the role of spinal cord oscillation, the appropriateness of the reference material properties for DCM, and the relationship between the measured stress and tissue injury. First, no studies specifically consider spinal cord oscillations [105]. The spinal cord oscillates cranio-caudally with heart rate. Recent imaging studies have indicated that this increased in the context of symptomatic stenosis, the nature of which may correlate with clinical measures of disease severity [106,107]. Spinal cord oscillation would likely result in a shear force on the spinal cord.

Second, it is uncertain how applicable the material properties are to DCM. Most elements are based on young healthy tissue references. In contrast, the ligaments and disks, for example, in DCM, are often degenerated and calcified, and, as aforementioned, the structure of the spinal cord is also recognized to change with age.

However, most importantly, none of these studies have specifically explored how the measured stress is related to tissue injury. Bridging this gap is critical, not only to fully confirm the appropriateness of FEA for DCM but also to guide its clinical interpretation [108]. All biological systems will have some baseline stress or strain; therefore, establishing disease thresholds will be critical to its development. The parallel development of in vivo techniques to measure tissue injury can complement this, for example, microstructural MRI and the less developed but promising serum and CSF biomarkers; however, this requires further prospective study.

Limitations
This study has some limitations. First, the search strategy focused on FEA models of the spinal cord and used citations to identify the source articles for all anatomical elements. Consequently, relevant source articles on the behavior of anatomical elements may have been missed. This is more likely for elements that were further removed from the spinal cord, such as the IVD, and experiments published more recently. This was a pragmatic decision based on the fact that existing investigators would likely have the best perspective on the literature, that this is a small research field, and that detailed biomechanical data on elements such as the IVD were unlikely to be so relevant. Consistent decisions across different research groups and findings across source articles would endorse this. Furthermore, due to the nature of our synthesis, we were unable to update our search. Although this may result in the omission of newer FEA articles, we believe that our review provides a useful approach for future investigators aiming to use FEA in DCM. Second, the methods used to shortlist source articles represent a framework we developed for the purpose of building a DCM FEA model. Again, the popularity of the shortlisted
articles across research groups provides some external validation, but it is possible that different investigators would reach different conclusions. For this reason, all source articles are listed in Multimedia Appendix 6, with their respective direct object identifiers. Third, this review aggregates data from a range of different experimental approaches and aims. Therefore, the analysis is largely qualitative, adhering to the SWiM guidelines [26]. Consequently, some conclusions, such as the relationship between the nature of spinal cord compression and stress distribution, remain tentative.

Conclusions
FEA has significant potential to help unlock uncertainties around the pathophysiology of DCM and inform clinical care. Currently, the application of FEA to DCM remains in its infancy. This review has adopted an intensive and iterative approach to help future investigators use FEA in DCM, including the aggregation of experimental data reporting on material properties and how they have been interpreted thus far. While single recommendations have been made, they have their limitations. The choice of material properties will influence the model performance, and investigators should consider their decisions carefully, particularly as new evidence emerges. More broadly, the methodology used in this review may be relevant to future updates and other clinical FEA initiatives when selecting material properties.

Acknowledgments
This study aligns with the AO Spine Research Objectives and Common Data Elements for Degenerative Cervical Myelopathy (RECODE-DCM) James Lind Alliance top research priorities, selected by people living and working with degenerative cervical myelopathy. This includes “biological basis,” and, to a lesser extent, “individualizing surgery” and “imaging and electrophysiology.” VN is supported by an NIHR Rosetrees Advanced Trust Fellowship. BMD was supported by a National Institute for Health Research Clinical Doctoral Research Fellowship. The views expressed in this publication are those of the authors and not necessarily those of the National Health Service, the National Institute for Health Research, Rosetrees Trust or the Department of Health and Social Care. The funders had no role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Data Availability
The data sets generated during and analyzed during this study are available from the corresponding author on reasonable request.

Authors’ Contributions
BMD designed the study, developed the search strategy, conducted the searches, screened the retrieved papers, extracted relevant information, and drafted the manuscript. SS contributed to paper screening and data extraction. ARF contributed to the writing of the subsequent drafts of this paper. VJFN and MPFS contributed throughout the project, starting from conceptualization to study design, search strategy development, and editing subsequent drafts of the paper.

Conflicts of Interest
None declared.

Multimedia Appendix 1
PRISMA-ScR (Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews) checklist.

Multimedia Appendix 2
Search strategy.

Multimedia Appendix 3
The quality assessment tool developed by the authors.

Multimedia Appendix 4
Comparison of modeling decisions.

Multimedia Appendix 5
Comparison of chosen equation and reference material property study.
References


Abbreviations

ACDF: anterior cervical discectomy and fusion
CSF: cerebrospinal fluid
DCM: degenerative cervical myelopathy
FEA: finite element analysis
IVD: intervertebral disk
MRI: magnetic resonance imaging
OPLL: ossification of the posterior longitudinal ligament
PRISMA-ScR: Preferred Reporting Items for Systematic Reviews and Meta-Analyses extension for Scoping Reviews
SCI: spinal cord injury
SWiM: Synthesis Without Meta-Analysis

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Sacroiliac Joint Dysfunction in Endurance Runners Using Wearable Technology as a Clinical Monitoring Tool: Systematic Review

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Abstract

Background: In recent years, researchers have delved into the relationship between the anatomy and biomechanics of sacroiliac joint (SIJ) pain and dysfunction in endurance runners to elucidate the connection between lower back pain and the SIJ. However, the majority of SIJ pain and dysfunction cases are diagnosed and managed through a traditional athlete-clinician arrangement, where the athlete must attend regular in-person clinical appointments with various allied health professionals. Wearable sensors (wearables) are increasingly serving as a clinical diagnostic tool to monitor an athlete’s day-to-day activities remotely, thus eliminating the necessity for in-person appointments. Nevertheless, the extent to which wearables are used in a remote setting to manage SIJ dysfunction in endurance runners remains uncertain.

Objective: This study aims to conduct a systematic review of the literature to enhance our understanding regarding the use of wearables in both in-person and remote settings for biomechanical-based rehabilitation in SIJ dysfunction among endurance runners. In addressing this issue, the overarching goal was to explore how wearables can contribute to the clinical diagnosis (before, during, and after) of SIJ dysfunction.

Methods: Three online databases, including PubMed, Scopus, and Google Scholar, were searched using various combinations of keywords. Initially, a total of 4097 articles were identified. After removing duplicates and screening articles based on inclusion and exclusion criteria, 45 articles were analyzed. Subsequently, 21 articles were included in this study. The quality of the investigation was assessed using the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) evidence-based minimum set of items for reporting in systematic reviews.

Results: Among the 21 studies included in this review, more than half of the investigations were literature reviews focusing on wearable sensors in the diagnosis and treatment of SIJ pain, wearable movement sensors for rehabilitation, or a combination of both for SIJ gait analysis in an intelligent health care setting. As many as 4 (19%) studies were case reports, and only 1 study could be classified as fully experimental. One paper was classified as being at the “pre” stage of SIJ dysfunction, while 6 (29%) were identified as being at the “at” stage of classification. Significantly fewer studies attempted to capture or classify actual SIJ injuries, and no study directly addressed the injury recovery stage.

Conclusions: SIJ dysfunction remains underdiagnosed and undertreated in endurance runners. Moreover, there is a lack of clear diagnostic or treatment pathways using wearables remotely, despite the availability of validated technology. Further research of higher quality is recommended to investigate SIJ dysfunction in endurance runners and explore the use of wearables for rehabilitation in remote settings.

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KEYWORDS
sacroiliac; sacroiliac dysfunction; sacroiliac wearables; sensors; injury management
Introduction

Physical activity, exercise, and sport are increasingly promoted as part of a healthy lifestyle. However, increased participation in physical activity and sport specialization may raise the risk of injury [1]. Running remains one of the most prevalent forms of physical activity, attracting individuals of all capability and ability levels to engage in this form of cardiovascular exercise. However, the burden of running-related injuries and their potential impact on quality of life and societal costs call for research and effective interventions in all the areas associated with sports injury, namely, prevention, assessment, and recovery [2,3]. One of the most overlooked sources of lower back pain (LBP) in endurance runners is injury to the sacroiliac joints (SIJs) [4].

The SIJs are the largest axial joints in the body and sit between the sacrum and pelvic bones on either side. The SIJs connect the spine to the pelvis and facilitate load transfer from the lumbar spine to the lower extremities. Specifically, the SIJs sit between the iliac’s articular surface and the sacral auricular surface. Therefore, the SIJ supports the torso and upper body muscular areas to dampen the impact of ambulation as the SIJ can experience forces of shearing, torsion, rotation, and tension when running. To improve and promote efficiency in running while focusing on injury prevention, allied health professionals are exploring different preventative, monitoring, and rehabilitative methods.

Numerous investigations have been undertaken to identify the factors contributing to the management of SIJ dysfunction and the underlying biomechanical mechanisms responsible for pain [3,4]. One consideration is using wearable sensor technology for clinical monitoring. In this regard, wearable sensors (wearables) incorporate a broad range of advances in microelectromechanical systems [5], electrocardiogram [6], electromyogram [7], and electroencephalogram-based neural sensing platforms [8]. As injuries such as SIJ dysfunction can require frequent monitoring, the continuousness of patient/athlete monitoring for timely intervention and rehabilitation seems essential. Wearables present an opportunity to measure the biomechanical parameters of SIJ dysfunction in a continuous, real-time, and nonintrusive manner by leveraging electronics packaging technology. It has been conveyed that by leveraging this technology, more time for engagement, continuity of experience, and dynamic data for decision-making for both athletes and clinicians will endure [9]. While remote and ambulatory monitoring are growing needs in the health care environment [10], the efficacy surrounding wearables in remote monitoring relative to SIJ dysfunction remains largely unknown. This is despite the acknowledgment that remote monitoring provides increased data volume and can promote improved athlete performance [11] and accelerate the patient/athlete rehabilitation processes [12]. Furthermore, an apparent limitation of existing research is that there has been a focus on the effectiveness of wearables on running performance metrics that generally do not consider ongoing rehabilitative considerations [13]. Strategies for the prevention of [14] and recovery from [3] SIJ injury have been proposed, alongside models of injury causation [15] and injury factors [16] (eg, intrinsic vs extrinsic; modifiable vs not modifiable). In turn, this has the potential to help monitor compliance, quality, and progress of movement performance when an injury-prevention or return-to-activity program is implemented [17]. Clinicians and allied health professionals often focus on exploring various training methods for preventive and rehabilitative measures. However, they rarely evaluate these methods in conjunction with biomechanical parameters and their impact on SIJ dysfunction. Thus, there is a need for evidence-based information on how wearables could be used for rehabilitation purposes in a remote setting when SIJ dysfunction is considered.

To maintain pace with the rapidly evolving field of wearables in endurance runners, this review provides an update on the state of the literature with a particular focus on literature published in the past 10 years. Case studies illustrate the use of wearable data in the development or monitoring of running programs. For the purposes of this review, a “wearable device” was operationally defined as a device that can be attached to the runner, shoe, or garment, or is a smartphone app. Thus, the purpose of this study was to systematically review the literature and gain a better understanding of the use of wearables in both in-person and remote settings for rehabilitation of SIJ dysfunction in endurance runners. Addressing this issue, the overall goal was to investigate how wearables can contribute to the clinical diagnosis (before, at, and after) of SIJ dysfunction.

Methods

Study Design

The design and reporting of this review followed the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses; Figure 1 and Multimedia Appendix 1 [18]) 2020 statement [18]. The general search strategy (Multimedia Appendix 2) and search terms are described in Table 1. Articles published up to October 1, 2022, were reviewed.

Thereafter, the selection process consisted of the following steps using the PRISMA guidelines (Figure 2): (1) an initial title screening for relevant articles was performed once the searched database results had been combined and duplicates had been removed; (2) both the titles and abstracts of the selected articles were then reviewed (a review of the full text was completed if it was not clear from the title or abstract whether the study met the review criteria); and (3) the full texts and selected articles were read based on the inclusion/exclusion criteria.
Figure 1. PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) flowchart.

Table 1. Systematic search strategy and key terms used.

<table>
<thead>
<tr>
<th>Search strategy</th>
<th>Key terms</th>
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<tbody>
<tr>
<td>Wearable technology</td>
<td>“Wearable Biomechanics” OR “Wearable Technology” OR “Wearable Devices” OR “Wearable Sensors Biomechanics” OR “IMU” OR “Inertial Sensor” OR “Inertial Measurement Unit” OR “Gyroscope” OR “Magnetometer” OR Accelerometer* OR “Pressure insoles” OR “Remote Wearables”</td>
</tr>
<tr>
<td>Running gait</td>
<td>“Running Biomechanics” OR “Endurance Running” OR “Run” OR “Jog” OR “Running over 5 km” OR “Endurance Runners” OR “Long Distance Runners” OR “Athletics”</td>
</tr>
<tr>
<td>Sacroiliac joint</td>
<td>“SIJ pain” OR “SIJ rehabilitation” OR “SIJ dysfunction” OR “SIJ injury prevention” or “SIJ management”</td>
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</table>

*TITLE-ABS-KEY was used as the search strategy.
A systematic search was conducted to identify potentially relevant papers in the following scientific databases: PubMed, Scopus, and Google Scholar. The focus of this review was on journal articles published in English that described the use of wearable technology to analyze, quantify, and emphasize the use of wearables for remote monitoring of SIJ dysfunction and rehabilitation in endurance runners. This extends to endurance runners undergoing rehabilitation for SIJ dysfunction (ie, had been diagnosed) or the ongoing management of SIJ dysfunction in previously diagnosed endurance runners (ie, rehabilitation). For this search strategy, an endurance runner was considered as someone partaking in regular running-related events (eg, recreational, fun runs) or competitive events (eg, competition, professional, elite). An endurance runner was classified as an athlete running more than 5 km in a single session, either during repeated trials or in studies that classified participants as endurance runners. In line with the main objective, inclusion and exclusion criteria were established to help eliminate studies that were not aligned with the research questions. An independent coder reviewed subsequent abstracts yielded from the search strategy and then the full articles for study selection. The review screened for information inclusive of health record and research systems including design, functionality, implementation, applications (remote and in-person settings) outcomes, and benefits. The search included articles published between 2000 and 2022. A manual review of the reference section of selected articles was then performed to identify relevant studies missed in the electronic search. Only English language articles were reviewed (Table 2).

**Inclusion and Exclusion Criteria**

A summary of the inclusion and exclusion criteria is presented in Table 2.

Although no restriction was imposed on the types of wearable technology used in SIJ dysfunction, the search terms were primarily focused on wearable inertial sensors and inertial measurement unit (IMU) devices (Table 3).
Table 2. Summary of inclusion and exclusion criteria.

<table>
<thead>
<tr>
<th>Study characteristic</th>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Communication type</td>
<td>• Journal and conference proceedings.</td>
<td>• Letters, short communications, technical notes, and other non–peer-reviewed literature. Non–evidence-based guidelines, letters to the editor, and expert opinion papers.</td>
</tr>
<tr>
<td>Injury classification</td>
<td>• Before, during, or after the clinical diagnosis of sacroiliac joint dysfunction which included or incorporated the use of wearables as a viable method of evaluating sacroiliac joint motion.</td>
<td>• Articles reporting exclusively on activity monitoring from global navigation satellite systems and injury surveillance without biomechanical measurements.</td>
</tr>
<tr>
<td>Classification of wearable</td>
<td>• Accelerometer, gyroscope, magnetometer, or a combination of these (inertial measurement unit), foot/shoe insoles (pressure mapping).</td>
<td>• Temperature sensors, pulse oximeters, pressure sensors, correlated glycemic measurement sensors, biosensitivity techniques, smartphone apps and related sensors, rehabilitation, and monitoring ambulator–based sensors.</td>
</tr>
<tr>
<td>Defined running gait outcome measure</td>
<td>• Spatiotemporal (global outcomes of the running gait cycle): running velocity, acceleration of the center of mass, distance, displacement, ground contact time, step length, step frequency (cadence), stance time, and flight time were included. Kinematics (description of segmental or joint movement, generally in the 3 cardinal planes, namely, sagittal, coronal [frontal], and transverse planes, without consideration for forces).</td>
<td>• Studies aiming to determine running power or economy were excluded as well as studies investigating walking gait variability or regularity. • Studies evaluating robotic systems, exoskeletons, prosthetics, and virtual reality environments were excluded. • Studies investigating the use of biofeedback or gait retraining (ie, nonnatural running gait) and studies involving the use of altered weight conditions (eg, wearable resistance, antigravity treadmills, or water-based protocols). • Computer algorithms; machine learning or statistical approaches; and those using robotic systems, exoskeletons, prosthetics, and virtual reality environments.</td>
</tr>
<tr>
<td>Kinetic</td>
<td>• Kinetic (the action of forces in producing or changing motion): for example, ground reaction force, peak pressure, center of pressure, braking, impulse, time to peak pressure, pressure time integral, loads, force time integral, and contact area.</td>
<td></td>
</tr>
<tr>
<td>Participant</td>
<td>• Age &gt;18 years, male and female. Endurance running included runners regularly completing over 5 km in training or competitive situations. • The endurance runner was partaking in regular running-related events (eg, recreational, fun runs) or competitive-based events (eg, competition, professional, elite). The runner was classified as an athlete running more than 5 km in a single protocol session, either during repeated trials or in studies that classified participants as endurance runners.</td>
<td>• Age &lt;18 years. Endurance runners not regularly completing over 5 km in training or competitive situations. • Studies done on animals and cadavers.</td>
</tr>
</tbody>
</table>
biomechanical screening and in-field injury events that injury occurrences in endurance running (e.g., cohort studies with mechanisms, and therefore, attempted to capture or track SIJ classifying SIJ injury factors, diagnosis, or underlying classified as the at category if they were identifying and SIJ dysfunction in endurance runners. Thus, studies were relationship between the experimental data collected and the “at/post” classification was used to express the chronological of SIJ dysfunction. Therefore, akin to Preatoni et al [20], an Studies were required to classify and characterize the diagnosis and characterize the diagnosis of SIJ dysfunction. Therefore, akin to Preatoni et al [20], an “at/post” classification was used to express the chronological relationship between the experimental data collected and the SIJ dysfunction in endurance runners. Thus, studies were classified as the at category if they were identifying and classifying SIJ injury factors, diagnosis, or underlying mechanisms, and therefore, attempted to capture or track SIJ injury occurrences in endurance running (e.g., cohort studies with biomechanical screening and in-field injury events that referenced use of wearable technology). Studies were classified as post if the data collection was performed after the SIJ injurious event, that is, during the SIJ recovery phase with the aim focused on rehabilitation techniques in both field-based and laboratory environments where the endurance runner had received a clinical diagnosis of SIJ dysfunction. The post classification was also used for studies that assessed the likelihood of SIJ injury or a greater magnitude of dysfunction. For clarity, studies that examined endurance runners who had returned to full running activity (e.g., comparisons between healthy individuals and those with a history of a specific or existing SIJ injury) were classified as pre because they were not centered on the recovery process that goes from injury occurrence (or medical intervention, if relevant) to being able to return to full running activity.

The characterization of studies was based on the following categories: (1) studies analyzing preexisting running-related SIJ dysfunction using wearable technologies to monitor running biomechanics in both a field-based and laboratory setting for the purpose of clinical management and clinical management in a remote setting (pre); (2) studies assessing endurance running–related SIJ dysfunction or injury factors or injury risk using wearable technologies to monitor running biomechanics after SIJ dysfunction has been formally diagnosed and classified as in the acute stage of injury in both field-based and laboratory settings for the purpose of clinical management at (at); (3) studies assessing ongoing running-related SIJ injury factors or injury risk using wearable technologies to monitor running biomechanics after SIJ dysfunction has been formally diagnosed and classified as in the chronic stage of injury in both field-based and laboratory settings for the purpose of clinical management or management in a remote setting (post); and (4) studies attempting to establish injury threshold criteria from a biomechanical perspective, studies characterizing protective wearable devices, and studies focusing on post-SIJ injury monitoring or return-to-run assessment using wearables. Validation and literature review studies were classified according to the primary aim for which the method or tool tested had been devised, as stated by the authors.

Study Classification and Assessment
The selected studies reported multiple feature domains, including (1) strength of evidence, time setting, and primary scope; (2) study characterization in terms of experimental conditions, setting (running field based and running laboratory based using treadmills), and age of endurance runners tested; and (3) characteristics of the technologies and types of wearable device and measures used relative to SIJ dysfunction. The author also defined and assessed (4) the Injury-research Readiness Level (IrRL) relative to SIJ dysfunction.

Selection Process: Strength of Evidence, Time Setting, and Scope
The strength of evidence for each article was assessed across 3 main categories, ordered in decreasing strength based on the experimental design used: experimental, that is, meeting the requirements of endurance running and SIJ dysfunction at or after clinical diagnosis and injury; randomized controlled trials; quasi-experimental, that is, including manipulation of the experimental conditions under which participants performed endurance running, but lacking random assignment or group comparison; and observational, that is, without assessing the effects of an intervention, and only describing participant behavior [19]. A separate class was used for studies looking exclusively at the validation of new equipment or methods. Literature reviews on wearables combined with synergies in remote settings or endurance running–related SIJ injuries were included and assessed by the primary author.

Classification and Characterization of SIJ Dysfunction
Studies were required to classify and characterize the diagnosis of SIJ dysfunction. Therefore, akin to Preatoni et al [20], an “at/post” classification was used to express the chronological relationship between the experimental data collected and the SIJ dysfunction in endurance runners. Thus, studies were classified as the at category if they were identifying and classifying SIJ injury factors, diagnosis, or underlying mechanisms, and therefore, attempted to capture or track SIJ injury occurrences in endurance running (e.g., cohort studies with biomechanical screening and in-field injury events that

Table 3. Comparative overview of wearable sensor modalities used in running today.

<table>
<thead>
<tr>
<th>Category</th>
<th>Technology</th>
<th>Capabilities</th>
<th>Pros</th>
<th>Cons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soft tissue injury prevention</td>
<td>Surface electromyogram</td>
<td>Identifies muscle recruitment and potential weaknesses</td>
<td>Small, wireless, and provides live data</td>
<td>Low signal-to-noise ratio</td>
</tr>
<tr>
<td>Workload management and</td>
<td>GPS, inertial measurement unit, and accelerometers</td>
<td>Distance, velocity, acceleration, deceleration, mediolateral movement, work, power, dehydration, fatigue, athletic performance, detecting gait parameters</td>
<td>Good range of data points</td>
<td>No biometric data and GPS can be pricey</td>
</tr>
<tr>
<td>athletic performance</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cardiac health</td>
<td>Electrocardiogram/photoplethysmography sensors</td>
<td>Heart rate, sleep rate, heart rate variability, respiration, muscle oxygen saturation, atrial fibrillation, stress levels, respiration rates, blood volume, and body temperature</td>
<td>Accurate and cost-effective</td>
<td>Price point</td>
</tr>
</tbody>
</table>

aThe table presents a comparative overview of common wearable sensors currently available rather than the components used for analysis (i.e., some studies used an inertial measurement unit, but only analyzed data from 1 element of the unit).

https://biomedeng.jmir.org/2024/1/e46067
To describe the experimental conditions, information was extracted about the settings in which data were collected (ie, laboratory vs field based). Specifically, studies were labeled as field based if wearable-obtained data were acquired during a scheduled running training event, a simulated running training event, or a running competition in a specific setting. Conversely, investigations carried out within a laboratory or in the field but using wearable technologies were labeled accordingly. The stage of SJJ dysfunction addressed by the study was then classified as either chronic (caused by overuse) or acute (resulting from specific events), following the criteria outlined by Bahr et al [15]. Furthermore, annotated classification of the endurance runner was addressed by each study (ie, recreationally active, trained/developmental, highly trained/national level, elite/international, world-class, or not specified/insufficient data to be classified) [21]. The risk of bias was assessed by the primary author.

**Injury-Research Readiness Level**

Building on the System Readiness Level framework by Sauser et al [22], an IrRL was modeled to capture the maturity, functionality, and readiness of the studies aiming to contribute to preventing, assessing, or recovering from SJJ dysfunction. According to the System Readiness Level model, technology and system development follow similar maturation paths, whereby technology is inserted into a system and interacts via a proposed architecture. Knowing about the system components and their integration is important, and this knowledge allows a classification of the system as being in its research, development, or deployment stage [23]. In the context of SJJ dysfunction and endurance running–related injuries, for this review, a method is deemed mature for deployment only when it relies on measuring wearable tools that are characterized by high ecological validity (ie, fully wearable and unobtrusive or markerless), can be applied directly in the field, is supported by validation studies against an established gold standard, or when validation is not practicable but adheres to standardized experimental procedures. Specifically, the biomechanical quantities pertaining to the SJJ should demonstrate evidence of a causal relationship with SJJ dysfunction and management in endurance running, and their interpretation should be driven by specific guidelines (eg, individual- or population-based normative boundaries, thresholds, or trends; Multimedia Appendix 3).

**Data Extraction and Collection**

After the data search was complete, data were obtained and extracted from eligible studies in a custom form that was created in Microsoft Excel. The form included (1) author, title, journal, and publication year; (2) research design; (3) sample size; (4) participant characteristics (eg, age, gender); (5) intervention features (type, length, and frequency); (6) measures and settings (laboratory, field-based, the type of wearable technology used, and sensors); (7) analysis; (8) key findings relative to the pre, at, and post categories for clinical SJJ dysfunction management using wearables in a remote or clinical setting; and (9) research outcomes, the metrics used, and conclusive statements. Data were then synthesized into a table format in Microsoft Excel and confirmed for data entry by the author. No automation tools were used in the process.

**Results**

**Overview of Identified Articles**

From the 4097 articles identified through the database search (Google Scholar, n=2263; Scopus, n=1624; and PubMed, n=210), and after removing duplicate items, 2245 publications were excluded based on title, abstract, and inappropriateness of topics (eg, knee arthroplasty in endurance runners). A further search was then performed in the databases with exclusion criteria (without the words) “knee” AND “lower back” AND “hip.” A search “with the words” was then refined to include “remote.” An additional 551 articles were removed due to “knee” appearing in the article while 2 papers were removed due to not being written in English. A further 4 were removed due to the topic being limited to physiological assessments only. A total of 1295 articles remained. Of these, 585 articles were discarded (most frequent reasons were not including wearables, not mentioning SJJ injury or SJJ dysfunction or running-related activities, and not describing the relationship between biomechanical quantities from wearables and the SJJ, or not defining the IrRL classification model relative to the SJJ and wearable usage in endurance runners). In addition, 665 records were removed due to technology not being classified as wearable, yielding a total of 45 studies to be considered for review.

A total of 151 participants were identified as being runners or endurance runners from the 45 papers analyzed. Descriptions of the included studies were either classified as a review of wearable sensors in the diagnosis and treatment of SJJ dysfunction, or wearable movement sensors for rehabilitation, or a combination of the above for SJJ gait analysis in an intelligent health care setting. Two papers [24,25] specifically mentioned wearable technology and the COVID-19 pandemic. However, only 1 of the review papers specifically mentioned measuring biomechanical loads and asymmetries in elite long-distance runners through inertial sensors [26]. One study [27] reported on SJJ pain relative to contralateral pelvic drop compared while the remaining research papers specifically mentioned iliac stress fractures in endurance runners linked to the SJJ, hip pain, or SJJ dysfunction. The remaining studies did not openly discuss the link between wearables and remote settings and SJJ dysfunction but mentioned such relationships as being possible or hypothetical. Thus, a total of 21 manuscripts remained, with overlapping reports on topics relative to SJJ dysfunction. No immediate forms of information bias (measurement bias) were detected in the final 21 studies.

**Journals and Years**

The 21 original manuscripts included in the review appeared in over 11 different journals, with 11 journals publishing nearly half of the total, and at least five relevant articles published in orthopedic, traumatology, or physical therapy journals. One paper was published in a rehabilitation journal while 3 papers were published in technology and engineering journals. The number of articles in the area under scrutiny appears to have increased over time, as 7 papers have been published since the
onset of the COVID-19 pandemic regarding telehealth (remote), sensors, and machine learning in endurance running injury management journals. The use of wearables in field-based and self-reliant monitoring seems to be increasing in popularity, as also demonstrated by the 7 review papers published between 2020 and November 2022.

More than half of the 21 studies scrutinized were literature reviews, 4 (19%) were case reports, and 1 was classified as fully experimental; 5 (24%) attempted to develop a predictive model or a machine learning approach to identify risk factors for running-related SIJ dysfunction. One study was classified as being at the pre stage of SIJ dysfunction, while 6 (29%) were identified as being at the at stage of classification. Considerably fewer studies attempted to capture or classify actual SIJ injuries, and no study directly addressed injury recovery (Table 4).
Table 4. Validity and reliability and application (information extracted from each article included the classification of the study).

<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Location</th>
<th>IrRL&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Classification</th>
<th>Participants and gender, n</th>
<th>Age (years)</th>
<th>Metric(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcantara et al [28]</td>
<td>2021</td>
<td>Force-measuring treadmill (laboratory)</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Validity (at&lt;sup&gt;b&lt;/sup&gt;)</td>
<td>37</td>
<td>Mean 20 (SD 2) years</td>
<td>Quantified accuracy of applying quantile regression forest and linear regression models to sacral-mounted accelerometer data to predict peak vertical ground reaction force, vertical impulse, and ground contact time across a range of running speeds.</td>
</tr>
<tr>
<td>Whitney et al [27]</td>
<td>2022</td>
<td>Treadmill (laboratory)</td>
<td>IrRL2: Development (building on established causal relationship)</td>
<td>Case-control (at)</td>
<td>81 runners (63 runners without SIJ pain and 18 runners with SIJ pain)</td>
<td>Mean 27.3 (SD 12.9) years for runners without and 23.8 (SD 10.5) years for runners with SIJ pain</td>
<td>In midstance, runners with SIJ pain had greater contralateral pelvic drop compared with controls. For unilateral SIJ pain cases (n=15), greater contralateral pelvic drop was observed when loading the affected side compared with the unaffected side. Female runners with SIJ pain demonstrated greater contralateral pelvic drop during the midstance phase, along with less knee flexion, greater “tibial overstride,” and greater ankle dorsiflexion at initial contact compared with controls.</td>
</tr>
<tr>
<td>Höfer and Siemsen [29]</td>
<td>2008</td>
<td>Treadmill (laboratory)</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Application (at) (proof of concept)</td>
<td>3 male participants</td>
<td>N/A</td>
<td>The pressure between the sensor contact area and the lumbar region was measured with force sensitive resistor sensors.</td>
</tr>
<tr>
<td>Ueberschär et al [26]</td>
<td>2019</td>
<td>Treadmill (laboratory)</td>
<td>IrRL2: Development (building on established causal relationship)</td>
<td>Experimental (pre&lt;sup&gt;c&lt;/sup&gt;)</td>
<td>45 healthy junior-elite long-distance runners</td>
<td>N/A</td>
<td>The mean peak tibial accelerations in junior-elite long-distance runners ranged between 14 (SD 3) and 16 (SD 3) g (g=9.81 m s&lt;sup&gt;-1&lt;/sup&gt;) for running speeds of 14–16 km h&lt;sup&gt;-1&lt;/sup&gt;. The corresponding mean peak sacral and scapular accelerations amounted to 4 (SD 1) to 5 (SD 1) g (32%, SD 8% of tibial load) and 4 (SD 1) g (mean 27%, SD 6%), respectively.</td>
</tr>
<tr>
<td>Liu et al [31]</td>
<td>2021</td>
<td>N/A</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Review</td>
<td>N/A</td>
<td>N/A</td>
<td>Daily monitoring of basic health data by wearable devices helps physicians in detecting the health problem. However, most current wearable sensors are not accurate enough for clinical evidence.</td>
</tr>
<tr>
<td>Banos et al [32]</td>
<td>2015</td>
<td>Laboratory</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Application (proof of concept/case report)</td>
<td>1 male participant</td>
<td>N/A</td>
<td>A novel mobile health system to support trunk endurance assessment. The system uses a wearable inertial sensor to track the patient’s trunk posture, while portable electromyography sensors were used to seamlessly measure the electrical activity produced by the trunk muscle.</td>
</tr>
</tbody>
</table>

<sup>a</sup> IrRL: Information reliability level

<sup>b</sup> at: Asymptomatic

<sup>c</sup> pre: Pilot study

<sup>d</sup> N/A: Not available
<table>
<thead>
<tr>
<th>Author</th>
<th>Year</th>
<th>Location</th>
<th>IrRL(^a) Classification</th>
<th>Participants and gender, n</th>
<th>Age (years)</th>
<th>Metric(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Falowski et al [33]</td>
<td>2020</td>
<td>N/A</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Review</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Zadeh et al [34]</td>
<td>2021</td>
<td>Laboratory</td>
<td>IrRL2: Development (building on established causal relationship)</td>
<td>Application (at) (proof of concept)</td>
<td>55 (39 male and 16 female participants)</td>
<td>21.1 (SD 3.84) years for male and 20.1 (SD 1.18) years for female participants</td>
</tr>
<tr>
<td>Porciuncula et al [35]</td>
<td>2018</td>
<td>N/A</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Review</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Lorusi et al [36]</td>
<td>2018</td>
<td>Field based</td>
<td>IrRL3: Deployment</td>
<td>Application (at) (proof of concept)</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Nascimento et al [37]</td>
<td>2020</td>
<td>N/A</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Review</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Rahlf et al [38]</td>
<td>2022</td>
<td>N/A</td>
<td>IrRL1: Research (exploring causal relationship)</td>
<td>Application (at) (proof of concept)</td>
<td>N/A</td>
<td>N/A</td>
</tr>
</tbody>
</table>

\(^a\)IrRL: Injury-research Readiness Level.

\(^b\)An at/post classification: if the scope was to identify and characterize SIJ injury factors, diagnosis, or underlying mechanisms; or track SIJ injury occurrences in endurance runners.

\(^c\)SIJ: sacroiliac joint.

\(^d\)N/A: not applicable.

\(^e\)Pre: pre-SIJ dysfunction (ie, before the SIJ injury).

**Experimental Setting**

In the field-based study [36] that analyzed endurance runners at the SIJ dysfunction stage, the application was at the proof-of-concept stage only. None of the studies included in this review were deemed to be experimental or classified as an observational study design pertaining to the use of wearables in a self-monitoring or remote rehabilitation capacity. This was despite most studies being literature or systematic reviews that focused on wearables for self-monitoring, self-monitoring in a remote setting, or a combination of both.

**Participant Characteristics**

Overall, the studies included between 1 participant [30] and 81 participants [26], with the mean number of participants being 21 (SD 32). The mean age of participants was 22.2 (SD 3.7) years. None of the selected studies performed a comparison of SIJ dysfunction and related gait patterns across the selected age groups or compared SIJ dysfunction using a validating approach in wearables. Many of the studies included both male and female participants; however, none of the selected studies examined differences between male and female participants in SIJ dysfunction using wearables. One study [26] focused on female runners with SIJ or sacral stress fractures, whereas another [29] included only male participants using pressure sensors in the lumbar region. Given the discrepancy in participant characteristics, a source of inequity, that is, gender bias, was prevalent in some studies analyzed.

**Clarification of SIJ Pathomechanics**

Overall, the SIJ appears to function as a stabilizer of the pelvis, absorbing ground reaction forces during gait and shear forces during movement [6]. The SIJ has also been described as a multidirectional force [39]. Activities that involve a 1-leg stance such as running would presumably increase the force in each SIJ, yet this was not specifically mentioned in the studies. Similarly, this would influence the vertical ground reaction force that occurs with each step. Another significant influence
is the center of mass, which is in slightly different positions for men and women. One study noted the importance of the center of mass, particularly in women, as it commonly passes in front of or through the SIJ [40]. Some of this can be explained due to sexual dimorphism being apparent in the pelvis, with the female sacrum being wider and with a more backward tilt. This would also account for the higher loads and stronger SIJs that are commonly seen in men [41]. This characteristic may also explain why men have more restricted mobility, as the average movement for men is approximately 40% less than that of women [42]. In this regard, the mechanism of SIJ dysfunction is primarily a result of a combination of axial loading and abrupt rotation [43]. DeRosa and Porterfield [44] delineated the primary influences as follows: the force of gravity, which acts downward through the spine, generating the flexion moment of the sacrum on the ilium, and the ground reaction force, which travels upward through the lower extremity from the heel strike, producing a posterior rotational moment (referred to as “torsional”) of the ilium on the sacrum; they termed these motions sacroiliac and iliosacral, respectively. Falowski et al [33] presented an algorithm for the diagnosis and treatment of SIJ pain. In this case, the authors believed that SIJ pain is an underdiagnosed and undertreated element of LBP. Citing an emerging disconnect between the growing incidence of diagnosed SIJ pathology and the underwhelming efficacy of medical treatment, they created a diagnostic and treatment pathway to establish an algorithm for patients that can include conservative measures and interventional techniques once the diagnosis is identified.

Classification of Wearables

A total of 8 studies used wearables in some form; however, only 1 study [26] used a sensor (a triaxial accelerometer) to measure biomechanical loads in endurance runners, although this study did not specifically review SIJ dysfunction. In the 8 studies that mentioned wearables, accelerometers and gyroscopes featured; however, the authors did not provide enough information to establish the type, range, and technical specification of the devices. There was a large variation in the reported use of temperature sensors, pulse oximeters, BioHarness wearable technology, pressure sensors, correlated glycemic measurements, biosensitivity techniques, electrodes, environmental monitoring, smartphone accelerometers, and next-generation wearable movement sensors despite these studies not specifically mentioning SIJ in endurance runners (Table 5).

Table 5. Breakdown of various approaches used for wearables.

<table>
<thead>
<tr>
<th>Approaches</th>
<th>Description</th>
</tr>
</thead>
</table>
| Referred to sensors’ validation within the cited article | • Compared with gold standards (eg, stereophotogrammetry, force platforms, high-speed video, or photocells) [24,25,34-37].  
• Comparing classification results against human/validated software classification [24,25,35-38]. |
| Pilot or proof studies                                | • Biomechanical effect of a lumbar spine-relief orthosis for the treatment of sacroiliac pain [29]. |
| Referred to ad hoc procedures for the performed measures | • Describing procedures for sacroiliac joint monitoring or pain management measures using machine learning or similar approaches [45,46]. |

The reviewed studies that used proof-of-concept designs [34,38] included generic descriptions of wearables relating to self-monitoring use and remote rehabilitation monitoring despite inadequate information provided about SIJ for rehabilitation in endurance runners. Furthermore, while describing the technical features of the wearable is key to the accurate clarification of data quality and of the implication of the changes that a remote intervention may encourage, many studies did not report this information sufficiently. Notably, and as highlighted by recent systematic reviews on wearables and inertial sensors for sport performance evaluation [47], and on accelerometer of impact loading in runners [30], reporting the features of the wearable device used—as well as information on the attachment location and fixing methods—is essential.

Discussion

Principal Findings

This review examined 21 studies that evaluated the effects of wearable use in remote settings during SIJ dysfunction in endurance runners. A secondary purpose of this review was to evaluate the effectiveness of wearables in possible or probable SIJ rehabilitation programs for endurance runners. Explicitly, this review reported on the (1) strength of evidence, time setting, and primary scope of studies relating to SIJ dysfunction in endurance runners; (2) characterization of SIJ dysfunction in terms of experimental conditions, setting (running field based or running laboratory based using treadmills), and the age of endurance runners tested; and (3) characteristics of the technologies and types of wearables and measures used relative to SIJ dysfunction in endurance runners. The author also defined and assessed (4) the IrRL relative to SIJ dysfunction. This review has demonstrated that the use of wearable technology for SIJ dysfunction monitoring in endurance running either from a laboratory or from a remote (telehealth) perspective is emerging, but further work is required to establish a standardized methodology and the validity or reliability of instrumentation.

This review provides a comprehensive overview of wearable technology used for an SIJ dysfunction in endurance runners as well as recommendations for future work.

Injury Type and Classification

The quality of the included studies varied, with one of the most challenging aspects of diagnosing and treating SIJ dysfunction...
in the endurance running population being the inconsistent judgment and, in some instances, worrisome presentation of the injury. The main difficulty faced by authors appears to be related to diagnostic challenges given that the pathomechanics and diagnostic classification of SIJ dysfunction are inconsistent in the literature. This was mainly observed in studies that referred to SIJ dysfunction as either a potential source of LBP or symbolic of hip-related issues. Moreover, lumbopelvic rhythm (LPR) was used as a definitive term by some authors. This, then, makes any possible deployment of wearables for rehabilitation purposes challenging if the diagnosis is either missed or misdiagnosed. As specific characteristics of SIJ dysfunction in endurance runners are required for investigation, the number of eligible participants was limited given that acute injuries were investigated primarily in 1 study [30] and chronic SIJ dysfunction in another [27], both of which occurred in control settings. None of the studies monitored acute or chronic SIJ dysfunction using wearables in a remote setting.

There were additional variations among the reviewed studies. While 2 studies examined the usability of wearables through active engagement with endurance runners [27,38], many lacked consideration for the wearer’s physical, psychological, and social preferences regarding the technology. Although 1 proof-of-concept study examined if wearable technology has the potential to predict injury in sports [34], many studies (42%) were found to be at the at stage of injury classification. However, it is important to consider the practicality of using wearables to classify SIJ dysfunction at the pre stage during running. Further research exploring the feasibility and necessity of using wearables is required, or whether this is feasible given the apparent difficulty in diagnosing SIJ dysfunction. Additional research will enhance our understanding of how wearables could be used at the onset of possible SIJ dysfunction to deliver the most pertinent data while enabling a clinical diagnosis.

A major issue in the approach to wearable instrument application is that more than half of the 21 studies analyzed were literature reviews, 4 (19%) were case reports, and 1 was classified as fully experimental relative to the classification of SIJ dysfunction. The results showed that although different wearables have been used for evaluating biomechanical parameters in the running gait analysis, as well as some relevant SIJ parameters pertaining to diagnostic or predictive stages of SIJ dysfunction, a paucity of research exists in the rehabilitation and remote monitoring of SIJ dysfunction. Indeed, the findings show that different descriptions related to possible or probable SIJ diagnosis exist in that injury classification is also referenced in relation to LBP and LPR. This, then, makes it difficult to draw firm conclusions regarding how wearables could be deployed remotely for rehabilitation purposes. Therefore, we are beginning to understand that the at stages of SIJ dysfunction require more than a concentration on the risk factors associated with injury occurrence.

Evidence also suggests that SIJ rehabilitation using wearable technology, in both controlled and remote settings, is highly nuanced (ie, varying across classification, injury stage, diagnosis, participant age, and gender). This complexity may extend to confusion in terminology and diagnosis between lower back injury and SIJ dysfunction, considering potential differences in running gait mechanics when running in controlled (eg, laboratory) versus remote settings. For example, one study [48] noted that the most common complaints were pain in the lower back, buttocks, leg, groin, and hip. Although some studies acknowledged that pain originating from the lower back region is likely more common than most endurance runners realize, as a result of the difficulty in localizing symptoms and referred pain patterns, the results suggest that reference to running-related SIJ issues was infrequent. This is not necessarily surprising as LBP is among the most common human health problems and accounts for a significant amount of disability worldwide [49]. Interestingly, the SIJ has been estimated to contribute to pain in as much as 38% of cases of LBP [50]. Although topographical classifications such as “sacroiliac,” “pelvis,” and “spine” serve a crucial didactic purpose, they can impede understanding of normal and altered functional SIJ mechanisms. As different classifications exist, it remains somewhat unknown if greater SIJ dysfunction in endurance runners exists, thus making any reference to the possible role of wearables relative to injury classification and rehabilitation in remote monitoring challenging.

What is commonly stated among the papers reviewed is that the clinical examination of an endurance runner with SIJ dysfunction commonly begins with an evaluation of gait. The results suggest that this often commenced in a clinical setting with ongoing monitoring of the condition commonly requiring the patient to be in the same clinical and controlled setting. It is at this juncture that wearables could be used in a remote and personalized setting, whereby data are fed to the clinician to monitor and track gait-related patterns or irregularities. Notwithstanding the literature reviews discussed in this paper that highlight the obvious and practical gap in using this technology in an SIJ dysfunction setting, more research is needed to test the feasibility and validity of the different wearable devices currently available. This extends to the level of expertise needed to operate and interpret the data from the perspective of an operator, athlete (runner) and clinician. Additionally, the results point to LPR being frequently referenced in the literature alongside LBP and SIJ dysfunction. The literature suggests that LPR is the relationship between the lumbar spine, hip, and pelvis when the trunk is in flexion. The classification of LPR during torso forward bending and backward return has also been widely investigated and commonly related to lower back disorders [51]. This defines LPR and LBP without necessarily drawing on the biomechanical differences and classification of how these injuries are managed in endurance runners. Furthermore, the results show considerable differences in the methods used to measure, and approaches used to characterize, LPR. Overall, it appears as though the timing aspect of LPR has been examined to obtain insights into the neuromuscular control of torso motion. The lack of consensus in LPR, LBP, and SIJ dysfunction is further impacted by the fact that there are no “gold-standard” algorithms for the detection of running gait outcomes from wearable sensor setups, which likely explains the large variation of outcomes and definitions reported in the reviewed studies.
Treatment of SIJ Dysfunction

It appears that treatment and management of SIJ dysfunction are often nonsurgical and involve packages of care that can include analgesics, physiotherapy, corticosteroid injections, and radiofrequency ablation [52]. Non-face-to-face (remote) care models exist in which the athlete is physically separated from the physician (or other health care workers) and empowered by communication-based technologies, such as videoconferencing and the use of continuous patient monitoring (wearable or “surface sensor”) technologies that capture athlete metrics and deliver health data remotely to the physician. Although these technologies have existed for some time, widespread implementation has been constrained by laws, regulations, and policies. The use of wearables in movement science and sport is widespread [53]; however, relative to SIJ dysfunction detection using wearables in either a laboratory/clinical setting or a remote setting, it could be argued that their application is still in an “exploratory phase.” Therefore, the findings agree with Hughes et al [54] in that the technology and the associated methods still require further development and careful analysis.

The results concerning SIJ injury risk mitigation have been well addressed in the literature [55,56]. Notwithstanding injury mitigation factors, no exploratory research has been performed to systematically investigate the feasibility of wearables use as a rehabilitation tool in SIJ injury assessment or dysfunction in endurance runners. This includes how wearables could potentially be used to characterize the severity of SIJ dysfunction as well as exploring the use of acquired information to support either clinical preventive or rehabilitative interventions. The empirical and analytical study of SIJ motion dates back to the late nineteenth century. However, its widespread acceptance as a legitimate entity has only occurred recently [57,58]. This delay in acknowledgment may elucidate why SIJ dysfunction can often be mistaken for LBP and LPR issues. Moreover, nowadays, the topography of SIJ motion should be measured to establish the conceivable axes of motion. From the study of Wilder et al [59], translation must occur for any sagittal innominate rotation to be possible because of the irregular surfaces and taut ligament structure. Accordingly, clinical theories have been proposed regarding the details of these motions. Along this line, Lee et al [6] stated that nutation seems to occur bilaterally when moving from supine to standing and unilaterally with flexion of the hip joint. Moreover, this kind of information would be relevant to any treatment of SIJ dysfunction given that counternutation occurs bilaterally and sometimes near the end of trunk flexion and unilaterally during hip extension. Some authors (eg, [60]) suggested that individuals with SIJ dysfunction display symmetrical gait and a depressed synergy between muscles providing SIJ force closure. The disorder involves reduced coactivation of the gluteus maximus and contralateral activation of the latissimus dorsi, which together provide joint stability during running. The disorder would be exacerbated in endurance runners given their need for maximum activation of gluteus maximus and torso stability, both of which require consideration when treating SIJ dysfunction. Nevertheless, these results indicate that the information on SIJ dysfunction in endurance runners and the treatment options that exist using wearables are unrepresented.

Despite these limitations, it is pertinent to consider whether such treatment methodologies are clinically and practically feasible within a given wearables context.

Information Technology and Health Care

Outcomes obtained from this review posit that health services have experienced great changes, especially in remote monitoring [61]. Additional clinical studies (eg, [31]) have shown that wearables are widely used to monitor functional and daily activity inclusive of walking and running gait. The wearables used were commonly integrated with an IMU sensor and controlled with a smartphone app [62]. The increased use of wearable technologies, either in isolation or as part of integrated, preventative, or rehabilitative approaches, offers an opportunity to collect quantitative data “in the field,” less obtrusively, for extended periods, and with fewer spatial limitations than conventional motion-capture technologies (eg, [46]). In this regard, wearables are increasingly viewed as promising alternatives to expensive analytical instruments in health care when specificity and selectivity criteria are met. It could be that wearables are used to monitor for possible pain, therefore exploring the use of torso acceleration as a proxy with a triaxial accelerometer. As the goals of SIJ dysfunction treatment may include increasing suppleness, strengthening, and correcting any asymmetries, the opportunity remains to explore how wearables could be used as a viable treatment monitoring option.

This, then, is an area for future research.

Wearables can help quantify spatiotemporal variables (eg, stride, step length, cadence) and physiology (eg, heart rate, recovery time) and are commonly used for human activity detection and quantified self-assessment. Until recently, or specifically since the emergence of the novel coronavirus, COVID-19 in January 2020, evidence for the effectiveness of remote usage and wearable monitoring, compared with traditional care models, has been scarce [63]. Along this line, the combination of telemedicine as an audiovisual communication platform and wearables that transmit field-based kinematic metrics provides numerous benefits to both health care providers and runners alike. Similarly, machine learning approaches have been widely used in gait biomechanics studies in the past decade [64-66]. However, among the papers included in this review, only 3 [31,32,36] focused on wearables for the sole purpose of remote monitoring of treatment of musculoskeletal disorders, clinical advances, and rehabilitation. This ambiguity further complicates the usage and uptake of wearables for SIJ dysfunction, which need to accommodate such conditions.

Although wearables can be used for home monitoring of activity and for the purposes of rehabilitation, little research has examined the potential of wearables when applied to acute or chronic SIJ dysfunction in endurance running. For example, when used remotely (ie, at home), the wearer (runner) could be required to complete standardized functional, rehabilitative assessments while data are continually recorded from the wearable device and relayed directly to the doctor or medic. Therefore, rather than comprising only standardized functional test data, as would be the case in a clinical setting, the runner’s ambulatory movement data set would contain data corresponding to all movements while wearing the sensor, including recovery.
and running activity. Indeed, common day-to-day movements can be tracked using wearable devices equipped with an IMU sensor and controlled with a smartphone app [62]. Besides research into wearable use in stride, step, stance, and spatiotemporal variables relative to both performance and injury mitigation, a greater understanding of the processes and predictors of SIJ rehabilitation has the potential to inform and strengthen public health. In this regard, the findings agree with Regterschot et al [67] in that important challenges and barriers to the deployment of wearables in clinical care remain. Similarly, Lang et al [68] discussed the major barriers to the application of wearables in motor rehabilitation and proposed benchmarks for the implementation of wearables in clinical practice. These clinical barriers include the demanding clinical environments that are often present, as well as the lack of recognition by some health professionals of the valuable information that can be obtained from wearables. There are also technology-related barriers, including (1) wearables that are inaccurate for many athletic populations (ie, inconsistent data output or lack of validity), (2) wearables that are not user-friendly for clinicians or athletes, and (3) the lack of published data on the reliability and clinical validity of some wearables. This extends to the development and optimization of innovative wearable configurations and data analysis techniques (eg, machine learning–based algorithms that enable the detection of specific activities and movements in free-living conditions). While Regterschot et al [67] asserted the existence of reliable and valid wearables for clinical populations and free-living environments, medical technology professionals could be encouraged to assist allied health specialists in developing the knowledge and skills necessary to effectively use wearables for remote rehabilitation purposes. In concordance with Regterschot et al [67], barriers exist in deploying remote wearables for detecting specific activities and movements in free-living conditions. The results of this review suggest that clinical barriers extend to the busy medical environment and the lack of realization of the value of information that can be obtained using wearables. However, it appears as though technological barriers also exist, including (1) a perception that wearables are inaccurate for many patient populations, (2) wearables that are not user-friendly for clinicians or patients, and (3) a lack of published data regarding reliability and clinical validity of sensor systems. Relatedly, Lang et al [68] discussed the clinical barriers to the application of wearables in motor rehabilitation and proposed benchmarks for the implementation of sensors in clinical practice. Therefore, researchers are encouraged to investigate the usability, acceptance, feasibility, reliability, and clinical validity of wearable sensors in clinical populations to facilitate the application of wearable movement sensors in SIJ rehabilitation.

Limitations

Some caution should be exercised when considering these findings. It merits noting that this review was a single-author systematic review. The author performed manual searches of all databases stated in this review and then coded and analyzed all retrieved results. Despite this, being a single-author review ensured that the processes described were based on the author’s judgment of eligible articles, albeit following the PRISMA guidelines diligently. While systematicity was adhered to as best as possible, a single-author review does incur a possible likelihood of unintentional bias and methodological limitations when compared with group reviews. However, the processes described by the author are based on data accumulation with clear links between the knowledge and content of the subject as well as providing evidence for future research. Additionally, this review is not meant to be exhaustive and includes only a cursory evaluation of the issues. The clinical applications discussed are limited to SIJ dysfunction in an endurance running population. As a potential limitation, endurance running was classified as involving runners regularly completing over 5 km in training or competitive situations. Therefore, papers featuring experimental trials involving runners covering distances below this threshold were not included. This was motivated by the very high publication rate that made their inclusion infeasible. Nevertheless, this potential limitation did not alter the key points raised in the large number of papers included in this review and presented in the Discussion section. While an effective SIJ is fundamental in one’s ability to run with biomechanical efficiency and effectiveness, this systematic review was not intended to review sensor-based methods solely for applied real-time gait analysis. As gait analysis can include sensors located on the shank and foot, which are most often used in combination with threshold or peak identification methods for gait detection for SIJ assessment, review papers on gait analysis were limited.

Recommendations

Despite these limitations, future studies should prioritize improving the quality of research aimed at reducing discrepancies in result interpretation, increasing reliability and validity, and promoting study generalizability. Given these findings, the review concurs with Block and Miller [69] that SIJ pain and dysfunction in endurance runners are likely highly underdiagnosed and undertreated. Additionally, clinicians should be mindful of a broader range of potential differential diagnoses regarding other sources of posterior hip and LBP in endurance runners.

Based on the findings of this review, wearables combined with smart devices could enable real-time data to be sent to health care professionals and clinicians, allowing for simultaneous tracking of endurance runners and monitoring the magnitude of SIJ dysfunction. This also challenges the engineering community to develop more intelligent, real-time, accurate information, making it user-friendly and offering athletes and clinicians actionable insights based on context-specific evaluation frameworks. As noted by Clermont et al [70], personalized and effective wearable technology should be rooted in a thorough understanding of the user’s experience, attitudes, and opinions which, if not properly considered, can severely hamper the potential of applications.

The selected articles, particularly those from 2020 and the onset of the COVID-19 pandemic, undoubtedly reflect the widespread interest in the area and an increasing trend in popularity. The analysis resulted in some key conclusions, which were reported along with main reflection points that led to the formulation of
guidelines and good practices for future research and dissemination. These are as follows:

- Articles should explicitly state the rationale for choosing and analyzing specific biomechanical quantities relating to the SIJ and include a justification of what relationship may exist between the SIJ and the diagnosed dysfunction. When previous literature and reviews are cited to support the choice made, the strength of evidence of previous studies should be discussed, together with the context from which that evidence emerged.
- More effort should be spent to fully exploit the potential of wearable technologies to detect and manage SIJ dysfunction, particularly as part of an injury management plan (post). This would allow the unobtrusive monitoring and quantification of the effects of prescribed interventions (preventive or rehabilitative) more regularly.
- The continuous progress in wearables offers many opportunities to collect data on many athletes simultaneously, unobtrusively, for long periods, and in field-based situations. However, the great “power” that even consumer-level technologies (eg, smartphones, watches, pods) currently offer does not come without problems, such as those associated with validity, user and clinician experience, and interpretation of data.

Conclusions

A current “state of play” in SIJ dysfunction among endurance runners for rehabilitation considerations using wearables in a remote setting was presented. This study took a systematic review approach to explore the existing literature on SIJ dysfunction in an endurance running population, using wearables as a rehabilitation tool. Viewed through the lens of wearable technology, the results from this review show that diagnosing, treating, and managing SIJ dysfunction in endurance runners vary considerably because of the inconsistent definition of the condition. To identify optimal rehabilitation considerations and effectively monitor this condition using remote wearables, further investigations are recommended to better clarify the condition. Moreover, greater utilization of wearables for measuring both biomechanics and pathomechanics is suggested to enhance the reliability and accuracy of remote wearable usage.

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Data Availability

The data sets used and analyzed during this study are available from the corresponding author upon reasonable request.

Conflicts of Interest

None declared.

Multimedia Appendix 1

PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) 2020 checklist. [DOCX File, 32 KB - biomedeng_v91e46067_app1.docx ]

Multimedia Appendix 2

Google Scholar, PubMed, and Scopus search strategies. [DOCX File, 21 KB - biomedeng_v91e46067_app2.docx ]

Multimedia Appendix 3

The IrRL classification model, where different classes of research maturity (IrRL1-3; columns) are mapped against the following feature domains (rows): knowledge of causal relationships, experimental settings, testing technology, and normative guidelines. IrRL: Injury-research Readiness Level. [DOCX File, 13 KB - biomedeng_v91e46067_app3.docx ]

References


https://biomedeng.jmir.org/2024/1/e46067

(JMIR Biomedical Engineering)


Abbreviations

<table>
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<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>IMU</td>
<td>inertial measurement unit</td>
</tr>
<tr>
<td>IrRL</td>
<td>Injury-research Readiness Level</td>
</tr>
<tr>
<td>LBP</td>
<td>lower back pain</td>
</tr>
<tr>
<td>LPR</td>
<td>lumbopelvic rhythm</td>
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</table>
**PRISMA**: Preferred Reporting Items for Systematic Reviews and Meta-Analyses

**SIJ**: sacroiliac joint

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Abstract

Background: The digital era has witnessed an escalating dependence on digital platforms for news and information, coupled with the advent of “deepfake” technology. Deepfakes, leveraging deep learning models on extensive data sets of voice recordings and images, pose substantial threats to media authenticity, potentially leading to unethical misuse such as impersonation and the dissemination of false information.

Objective: To counteract this challenge, this study aims to introduce the concept of innate biological processes to discern between authentic human voices and cloned voices. We propose that the presence or absence of certain perceptual features, such as pauses in speech, can effectively distinguish between cloned and authentic audio.

Methods: A total of 49 adult participants representing diverse ethnic backgrounds and accents were recruited. Each participant contributed voice samples for the training of up to 3 distinct voice cloning text-to-speech models and 3 control paragraphs. Subsequently, the cloning models generated synthetic versions of the control paragraphs, resulting in a data set consisting of up to 9 cloned audio samples and 3 control samples per participant. We analyzed the speech pauses caused by biological actions such as respiration, swallowing, and cognitive processes. Five audio features corresponding to speech pause profiles were calculated. Differences between authentic and cloned audio for these features were assessed, and 5 classical machine learning algorithms were implemented using these features to create a prediction model. The generalization capability of the optimal model was evaluated through testing on unseen data, incorporating a model-naive generator, a model-naive paragraph, and model-naive participants.

Results: Cloned audio exhibited significantly increased time between pauses (P<.001), decreased variation in speech segment length (P=.003), increased overall proportion of time speaking (P=.04), and decreased rates of micro- and macropauses in speech (both P=.01). Five machine learning models were implemented using these features, with the AdaBoost model demonstrating the highest performance, achieving a 5-fold cross-validation balanced accuracy of 0.81 (SD 0.05). Other models included support vector machine (balanced accuracy 0.79, SD 0.03), random forest (balanced accuracy 0.78, SD 0.04), logistic regression, and decision tree (balanced accuracies 0.76, SD 0.10 and 0.72, SD 0.06). When evaluating the optimal AdaBoost model, it achieved an overall test accuracy of 0.79 when predicting unseen data.

Conclusions: The incorporation of perceptual, biological features into machine learning models demonstrates promising results in distinguishing between authentic human voices and cloned audio.

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KEYWORDS

voice; vocal biomarkers; deepfakes; artificial intelligence; vocal; sound; sounds; speech; audio; deepfake; cloning; text to speech; cloned; deep learning; machine learning; model-naive
Introduction

An increasing number of individuals rely on digital platforms as their primary sources of news and information [1]. People often trust what they consume on the internet without doing any research on the source. There is a technological advancement significantly influencing the production of digital media known as “deepfake.” Deepfake constitutes a synthetic reproduction of media content, both auditory and visual, carefully crafted to closely represent the physical attributes and vocal characteristics of a specific individual. Its use spans many domains, notably in entertainment, where it can be used for the digital replication of actors for special effects or the creation of intricately detailed characters in video games [2].

Deepfakes are generated through the aggregation of substantial data sets, including voice recordings, images, and video segments [3]. This research specifically targets the detection of audio deepfakes, relying solely on voice data for both deepfake development and detection method testing. The voice data sets serve as the foundation for training deep learning models, predominantly deep neural networks, with the primary objective of encoding unique and distinguishable attributes and characteristics found in human voices, like speech patterns and intonation [3]. Following successful model training, it gains the capability to produce replicated voice data by processing input audio or text [3]. While initially trained with substantial data sets, deepfake generation models posttraining can produce new voice clones with minimal audio input, synthesizing voice data to replicate the target voice’s distinctive traits based on learned patterns during the training phase.

This technology is valuable in many domains including voice assistants, voice dubbing for multimedia, professional voiceovers, and the narration of audiobooks [4]. Deepfake content can be generated rapidly once a model is trained, thereby significantly improving efficiency across many industries. Unfortunately, the irresponsible and unethical misuse of deepfakes is prevalent, encompassing impersonation, the dissemination of false information, and violation of privacy [5,6]. Due to the dynamic and rapidly evolving nature of this technology, remaining updated with the ongoing advancements in deepfake detection is challenging [7].

Individuals need a reliable tool to verify that the information they are consuming is authentic. Several outdated deepfake detection machine learning methods have high levels of accuracy, achieving up to 100% accuracy on a data set [8]. However, these accurate predictions are restricted to the level of advancement of the deepfakes that the detection models are trained with [9]. For example, the previously mentioned tool that achieved 100% accuracy was trained and tested on a data set of deepfakes generated in 2019, which are of much lower quality than the level of deepfakes available in 2023 [8]. Furthermore, recent work has shown that out-of-domain voice clone detectors (ie, voice detectors applied outside of the data set in which they were applied) had extremely low performance, obtaining an area under the receiver operator curve (AUC) of 25% [10]. A more robust detection method might involve searching for the absence of biological features in the cloned voice, rather than the presence of digital features [11].

Activities such as respiration, swallowing, and cognitive processes can influence speech production and the pattern of pauses in authentic speech. Although voice cloning processes may closely mimic human speech production, machines have no requirements for speech breaks and instead rely on training data to indicate where these pauses occur. This may result in subtle but detectable differences in the way pauses are present in authentic versus cloned audio. Indeed, when humans were asked to distinguish between audio deepfakes and authentic voices, one of the primary justifications for a fake audio classification was unnatural pauses in the recordings [10]. Furthermore, when these features were integrated into a classification regime, a moderate accuracy (approximately 85%) was achieved when analyzing deepfakes by perceptual features such as the amplitude of speech and pauses within a recording [12]. However, that study only assessed the use of a single voice cloning software (ElevenLabs) and a small number of cloned voices (9 built-in text-to-speech (TTS) voices and voices cloned from 2 celebrities). Furthermore, the training, validation, and testing sets were not split by participants, so it is assumed that recordings from the same participant are present in both the training and testing data sets.

We posit that the absence of regular human vocal biomarkers, characterized by the pause pattern in a speech segment, will be effective in differentiating cloned audio from authentic audio. For a more comprehensive understanding of model performance on out-of-domain data, we test the proposed methodology in the following ways:

1. On real and cloned audio recordings the model was not exposed to during training, including built-in TTS obtained from the cloning models
2. On a paragraph the model was not exposed to during training
3. On a new cloning software the model was not exposed to during training

Methods

Recruitment

A total of 49 adult participants (20 male) were recruited for this study between June and August 2023 in Toronto, Canada. The participant pool exhibited diversity in terms of ethnicity and had various types and strengths of accents. Exclusion criteria for recruitment included: (1) any person not living in Canada, (2) any person below the age of 18 years, and (3) any speech pathology or condition impeding the production of standard speech, such as stuttering, vocal cord pathology, tracheostomy, or the common cold. No restrictions on gender, ethnicity, accents, or other demographic data were implemented in the recruitment procedure.

The summarized protocol, as illustrated in Figure 1, involves participants recording the required voice samples for the training of 3 distinct deepfake models and a control version of 3 test paragraphs. Subsequently, each deepfake model generates each test paragraph, resulting in a total of 9 deepfake audio samples,
in addition to the 3 control samples for each participant. It is worth noting that some participants were unable to complete the necessary training voice recordings for 1 or 2 of the deepfake generators due to time constraints, resulting in varying numbers of recordings and deepfakes among participants.

**Figure 1.** General study protocol overview comprising the audio collection section and detection model development for a participant used in model training. Note that for participants not used in model training (“Model-Naïve Participants”), all data are used for model testing. ADA: AdaBoost; DT: decision tree; LR: logistic regression; RF: random forest; SVM: support vector machine.

**Ethics Approval**

The research protocol received approval from the Canadian SHIELD Ethics Review Board (REB Tracking Number 2023-06-003).

**Audio Samples**

In this study, we generated deepfakes using 3 publicly available and user-friendly web-based models: ElevenLabs [13], Podcastle [14], and Descript [15]. Each of these models required different training data. ElevenLabs had the least specific training requirements and was provided approximately 10 minutes of voice recordings, Descript required 10 minutes of speech samples, and Podcastle required participants to read 70 short phrases.

Recordings took place in a quiet room with participants seated in front of a MacBook Pro with 2.8 GHz Quad-Core Intel Core i7. They were instructed to articulate their speech clearly at a standard speaking volume, using the laptop’s built-in microphone to record. The laptop screen displayed the text that participants were required to read for the collection of voice sample data, including the 3 test paragraphs used in the development of the classification model.

All audio samples were saved in the Waveform Audio Format. The respective voice sample data were input for each deepfake generation model for the training process. Upon completion of the model training, a TTS technique was used to generate deepfake versions of the 3 test paragraphs for each model.

Each voice cloning platform also provides pregenerated TTS voices. We generated each of the 3 paragraphs using all available pregenerated TTS to be used in model testing.

**Feature Generation**

The aim of the analysis was to characterize cloned voices using amplitude-agnostic perceptual voice features, primarily characterized by the pause patterns within a speech segment. Speech segments were identified using a voice activity detector (VAD Solero) in Python [16]. The time between speech segments was calculated and classified as a micropause if the time between segments was greater than or equal to 0.1 seconds and less than 0.5 seconds. It was classified as a macropause if the time between segments was greater than or equal to 0.5 seconds (Figure 2). The recording was trimmed so that the recording began at the beginning of the first speech segment and concluded at the end of the final speech segment. Overall, five features were obtained to denote the pause pattern:
1. SpeechAV: The average speech segment length.
2. SpeechSD: The SD of the speech segment lengths.
3. SpeechProp: The proportion of time speaking, calculated by the sum of all the speech segment lengths divided by the length of the entire recording.
4. MiRate: The rate of micropauses, calculated by dividing the number of micropauses by the length of the trimmed recording (in minutes).
5. MaRate: The rate of macropauses, calculated by dividing the number of macropauses by the length of the trimmed recording (in minutes).

Previous work published by Barrington et al [12] evaluated perceptual features to compare audio deepfakes and authentic voices. In this work, 4 summary metrics to characterize the pauses were generated: the average length of a pause, the SD of the pauses, the pause ratio, and the total number of pauses. We slightly modified and expanded these features to align with our hypothesis. Rather than the average length and SD of the pauses, we used the average length and SD of the speech segments. We hypothesized that cloned audio would have longer periods between pauses, as they would have no requirements for biological processes such as breathing or swallowing. Furthermore, instead of reporting the number of pauses, which is dependent on the text spoken and the length of the recording, we exclusively reported pause rates. To account for the differences in pause lengths, we calculated the rates of both micropauses and macropauses.

Contrary to the work published by Barrington et al [12], we chose not to include amplitude features. The amplitude of a voice recording can be influenced by the type of microphone used in recording and the distance of the participant to the microphone. Due to this variation, and the desire to evaluate pause metrics exclusively, we chose to remove amplitude-associated features from our feature set.

Figure 2. Sample speech and pause illustration. Black segments indicate speech segments, red segments illustrate micro pauses (pauses $<0.5$ seconds and $\geq 0.1$ seconds), and yellow segments indicate macro pauses (pauses $\geq 0.5$ seconds).

Audio Feature Information
Audio features were compared between authentic and cloned audio. All analysis was conducted in Python. Statistical analysis was conducted using the scipy Python package [17]. P values were calculated using the Mann–Whitney U test. Statistical significance is defined as $P<.05$.

Detection Model Generation
An experiment was conducted to assess 5 models to determine the most suitable machine learning tool for this application: random forest (RF), decision tree (DT), logistic regression (LR), support vector machine (SVM), and AdaBoost (ADA) models. Neural networks, although useful in previous deepfake detection methods, perform best with large amounts of training data and tend to overfit with smaller data sets. We aimed to show speech pause patterns could be used to create a robust model even with a small amount of training data, so neural networks were not included in the current analysis.

A 5-fold stratified group cross-validation was used during model training and hyperparameter tuning to find the optimal model. Paragraphs 1 and 2 in Multimedia Appendix 1, and ElevenLabs and Podcastle generators were used in model training. A total of 30 participants were used in cross-validation (approximately 60% of participants). All recordings corresponding to a participant were kept in the same group, such that if a participant was in one of the folds, all the authentic and cloned recordings obtained from that participant were in the same fold. The total number of recordings used in cross-validation model training is displayed in Table 1.

All analysis was conducted in Python. Models were trained using the scikit-learn Python package [18]. Hyperparameters were tuned using the GridSearch algorithm in scikit-learn, using the parameters denoted in Multimedia Appendix 2. Accuracy is defined as

Model performance was assessed by the average balanced accuracy of all folds for a model, defined as
where \( k \) is the fold number, sensitivity is the accuracy of the model in predicting audio deepfakes, and specificity is the accuracy of the model in predicting authentic audio.

### Table 1. Number of recordings collected and generated.

<table>
<thead>
<tr>
<th></th>
<th>Training data set (P1\textsuperscript{a}/P2\textsuperscript{b}), n</th>
<th>Testing data set (P1/P2/P3\textsuperscript{c}), n</th>
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<td>257 (63/58/136)</td>
<td>384 (126/122/136)</td>
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<td>ElevenLabs</td>
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<td></td>
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<td>Pretrained</td>
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<td>19 (7/5/7)</td>
<td>19 (7/5/7)</td>
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<td>83 (24/22/37)</td>
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<tr>
<td>Cloned</td>
<td>—</td>
<td>46 (13/16/17)</td>
<td>46 (13/16/17)</td>
</tr>
<tr>
<td>Total recordings</td>
<td>—</td>
<td>52 (15/17/18)</td>
<td>52 (15/17/18)</td>
</tr>
<tr>
<td>Authentic</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total recordings</td>
<td>55 (28/27)</td>
<td>75 (13/13/49)</td>
<td>130 (41/40/49)</td>
</tr>
</tbody>
</table>

\( \textsuperscript{a} \)P1: paragraph 1.  
\( \textsuperscript{b} \)P2: paragraph 2.  
\( \textsuperscript{c} \)P3: paragraph 3.  
\( \textsuperscript{d} \)Not applicable.

### Optimal Model Testing

The optimal model from the detection model generation was tested on unseen data. For testing, there were three subgroups of data:

1. Audio recordings from individuals the model was not exposed to during training. This subgroup consists of:
   - Participant audio recordings that were not used in model training ("Model-Naïve Participants"). Note that for a participant to be “Model-Naïve”, neither authentic nor cloned audio obtained from that participant was used in model training.
   - Built-in, pretrained TTS obtained from the cloning models ("Pre-Generated TTS")

2. A paragraph the model was not exposed to during training ("Model-Naïve Paragraph": P3, Multimedia Appendix 1).

3. A new cloning software the model was not exposed to during training ("Model-Naïve Generator"). This was the Descript generator.

The model was tested in such a way that each testing datapoint was Model-Naïve in at least 1 of the 3 above subgroups. Data classes used in model training are denoted as “Model-Trained”.

### Results

#### Audio Feature Information

The 5 audio features corresponding to the speech pause profiles were calculated from the training data and are displayed in Table 2. Overall, cloned audio was significantly associated with increased time between pauses (\( P<.001 \)), decreased variation in the length of speech segments (\( P=.003 \)), increased overall proportion of time speaking (\( P=.04 \)), and a decreased rate of micro- and macropauses in speech (both \( P=.01 \)).
Table 2. Participant and recording data for model features for training data.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Authentic audio, mean (SD)</th>
<th>Cloned audio, mean (SD)</th>
<th>P valuesa</th>
</tr>
</thead>
<tbody>
<tr>
<td>SpeechAV</td>
<td>2.93 (1.76)</td>
<td>3.49 (1.23)</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>SpeechSD</td>
<td>1.51 (1.83)</td>
<td>1.22 (0.89)</td>
<td>.003</td>
</tr>
<tr>
<td>SpeechProp</td>
<td>0.87 (0.04)</td>
<td>0.89 (0.04)</td>
<td>.04</td>
</tr>
<tr>
<td>MiRate</td>
<td>11.72 (4.34)</td>
<td>9.47 (4.25)</td>
<td>.01</td>
</tr>
<tr>
<td>MaRate</td>
<td>7.04 (3.39)</td>
<td>5.78 (2.74)</td>
<td>.01</td>
</tr>
</tbody>
</table>

aP value calculated using Mann-Whitney U test. Statistical significance defined as P<.05.

Detection Model Generation

Five classical machine learning algorithms were implemented to create the prediction model, using the 5 features presented in Table 2. A total of 127 recordings were used to train each model and 257 recordings were used to test each model (see Table 1). The optimal performance was obtained by an ADA model, achieving a 5-fold cross-validation balanced accuracy of 0.81 (SD 0.05). The subsequent models were SVM (balanced accuracy 0.79, SD 0.03) and RF (balanced accuracy 0.78, SD 0.04), followed by LR and DT (balanced accuracies 0.76, SD 0.10 and 0.72, SD 0.06). Unsurprisingly, the models that are traditionally less prone to overfitting (ADA and SVM) were the models that had the best performance, whereas the model that was more likely to overfit (DT) had the poorest performance. Furthermore, ADA and other boosted models can experience the curse of dimensionality when data have many features. By using a small feature set (5 features), we avoided this problem, and ADA achieved a high cross-validated accuracy. Receiver operator curves of all models are shown in Figure 3, and additional model metrics are presented in Table 3. Tuned model hyperparameters are presented in Multimedia Appendix 2.

Figure 3. Average receiver operator curves with variability of all models. The results presented are calculated using the optimal parameter set for each model after Grid Search cross-validation. ADA: AdaBoost; AUC: area under the receiver operator curve; DT: decision tree; LR: logistic regression; RF: random forest; ROC: receiver operator curves; SVM: support vector machine.
Table 3. Model prediction results for all models.

<table>
<thead>
<tr>
<th>Model</th>
<th>Balanced accuracy, mean (SD)</th>
<th>Authentic voice accuracy, mean (SD)</th>
<th>Cloned voice accuracy, mean (SD)</th>
<th>Precision, mean (SD)</th>
<th>F1-score, mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AdaBoost^b</td>
<td>0.81 (0.05)</td>
<td>0.75 (0.09)</td>
<td>0.87 (0.08)</td>
<td>0.82 (0.07)</td>
<td>0.84 (0.04)</td>
</tr>
<tr>
<td>Support vector machine</td>
<td>0.79 (0.03)</td>
<td>0.73 (0.06)</td>
<td>0.85 (0.05)</td>
<td>0.80 (0.03)</td>
<td>0.82 (0.02)</td>
</tr>
<tr>
<td>Random Forest</td>
<td>0.78 (0.04)</td>
<td>0.73 (0.08)</td>
<td>0.83 (0.07)</td>
<td>0.80 (0.07)</td>
<td>0.81 (0.05)</td>
</tr>
<tr>
<td>Logistic Regression</td>
<td>0.76 (0.10)</td>
<td>0.70 (0.16)</td>
<td>0.83 (0.09)</td>
<td>0.79 (0.11)</td>
<td>0.81 (0.08)</td>
</tr>
<tr>
<td>Decision Tree</td>
<td>0.72 (0.06)</td>
<td>0.71 (0.08)</td>
<td>0.73 (0.15)</td>
<td>0.77 (0.07)</td>
<td>0.73 (0.09)</td>
</tr>
</tbody>
</table>

^aResults presented are calculated using the optimal parameter set for each model after Grid Search cross-validation.
^bOptimal model.

Optimal Model Testing
The optimal ADA model was tested on trained and naïve generators and participants with the paragraphs used in model training (Table 4), and a Model-Naïve paragraph (Table 5). The optimal overall testing performance was obtained when the model was tested on pretrained paragraphs for naïve participants (0.89 overall accuracy). The poorest authentic classification accuracy was obtained when trained participants spoke a new paragraph (accuracy 0.70), potentially indicating the model was overfit to the paragraphs used in training by trained participants. The highest authentic classification accuracy was obtained by model-naive participants speaking model-trained paragraphs with an accuracy of 0.96. Conversely, the detection of cloned and pregenerated voices typically performed better on Model-Naïve paragraphs (most accuracies >0.70). The exception to this was the Model-Naïve Generator which had an overall accuracy of 0.67. However, the number of datapoints for this category was extremely small (N=3) so this accuracy may not be the best representation of the Model-Naïve Generator performance. Pregenerated voices with the trained paragraphs had the lowest performance of all the model testing (overall 0.67 accuracy), but classification performance was much higher in the model-naive paragraph (overall accuracy 0.89). When the results of all confusion matrices in Tables 4 and 5 are compiled, the overall accuracy of all testing data was 0.79 with an AUC of 0.88.

Table 4. Confusion matrices of model test results for model-trained paragraphs (P1 and P2).

<table>
<thead>
<tr>
<th></th>
<th>Predicted authentic</th>
<th>Predicted fake</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model-trained participants</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Authentic</td>
<td>—^a</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Model-trained generator</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Model-naïve generator</td>
<td>5</td>
<td>17</td>
<td>0.773</td>
</tr>
<tr>
<td>Overall</td>
<td>—</td>
<td>—</td>
<td>0.773</td>
</tr>
<tr>
<td>Model-naïve participants</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Authentic</td>
<td>25</td>
<td>1</td>
<td>0.962</td>
</tr>
<tr>
<td>Model-trained generator</td>
<td>3</td>
<td>10</td>
<td>0.769</td>
</tr>
<tr>
<td>Model-naïve generator</td>
<td>1</td>
<td>7</td>
<td>0.875</td>
</tr>
<tr>
<td>Overall</td>
<td>—</td>
<td>—</td>
<td>0.894</td>
</tr>
<tr>
<td>Pregenerated TTS^b</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Authentic</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Model-trained generator</td>
<td>17</td>
<td>31</td>
<td>0.646</td>
</tr>
<tr>
<td>Model-naïve generator</td>
<td>0</td>
<td>4</td>
<td>1.00</td>
</tr>
<tr>
<td>Overall</td>
<td>—</td>
<td>—</td>
<td>0.673</td>
</tr>
</tbody>
</table>

^aNot applicable.
^bTTS: text-to-speech.
Table 5. Confusion matrices of model test results for the Model-Naïve paragraph (P3).

<table>
<thead>
<tr>
<th>Model-trained participants</th>
<th>Predicted authentic</th>
<th>Predicted fake</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Authentic</td>
<td>19</td>
<td>8</td>
<td>0.704</td>
</tr>
<tr>
<td>Model-trained generator</td>
<td>7</td>
<td>29</td>
<td>0.806</td>
</tr>
<tr>
<td>Model-naïve generator</td>
<td>0</td>
<td>14</td>
<td>1.00</td>
</tr>
<tr>
<td>Overall</td>
<td>—</td>
<td>—</td>
<td>0.805</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Model-naïve participants</th>
<th>Predicted authentic</th>
<th>Predicted fake</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Authentic</td>
<td>16</td>
<td>6</td>
<td>0.727</td>
</tr>
<tr>
<td>Model-trained generator</td>
<td>1</td>
<td>7</td>
<td>0.875</td>
</tr>
<tr>
<td>Model-naïve generator</td>
<td>1</td>
<td>2</td>
<td>0.667</td>
</tr>
<tr>
<td>Overall</td>
<td>—</td>
<td>—</td>
<td>0.758</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Pregenerated TTS\textsuperscript{b}</th>
<th>Predicted authentic</th>
<th>Predicted fake</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Authentic</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Model-trained generator</td>
<td>3</td>
<td>21</td>
<td>0.875</td>
</tr>
<tr>
<td>Model-naïve generator</td>
<td>0</td>
<td>2</td>
<td>1.00</td>
</tr>
<tr>
<td>Overall</td>
<td>—</td>
<td>—</td>
<td>0.885</td>
</tr>
</tbody>
</table>

\*Not applicable.

\textsuperscript{b}TTS: text-to-speech.

Discussion

### Principal Findings

This paper outlines the development of an audio deepfake detection model that capitalizes on the distinctive biological vocal characteristics to distinguish between genuine human speech and machine-generated audio. Voice clone samples were created for each participant using 3 publicly available platforms: Descript, ElevenLabs, and Podcastle. To compare these cloned samples with the participants’ authentic voice recordings, a variety of perceptual features were calculated to characterize the pause pattern in a recording. The hypothesis was that the speech and pause pattern would be distinguishable between authentic voice recordings and voice clones, as a machine-generated audio sample would not be under the same biological requirements as a human. Machines have no requirements for breathing or swallowing, and their processing time is magnitudes shorter than humans. Even if machines falsely replicate the pauses in speech, their lack of necessity for these processes may create subtle distinctions in the overall pause patterns. Our results support this finding, and 5 perceptual pause features were used to create a detection model for cloned audio.

To generate the voice classification model, 5 machine learning algorithms were used. An ADA model emerged as the most capable of classification, achieving an accuracy of 0.81 (SD 0.05) in 5-fold cross-validation and similar accuracy (0.79) across all testing experiments. The accuracy is in line with previous pause rate detection methods [12], although the testing methodology presented here allows for more comprehensive conclusions about the extendibility of the model results and possible implications for future work. Overall, Model-Naïve participants, a variety of generators, and Model-Naïve paragraphs were used to test the feasibility of the approach.

In the 5-fold cross-validation model optimization, we achieved an accuracy of 0.75 (SD 0.09) for authentic audio and 0.87 (SD 0.08) for cloned audio. Authentic accuracy may have been lower due to the inherent variation in real human speech, as demonstrated by the higher SDs of the pause metrics in Table 1 compared with cloned audio. This could result in decreased performance, as authentic audio may be more likely to overlap with cloned audio features and thus be harder to classify. Furthermore, we did not prioritize authentic speech accuracy in cross-validation, instead optimizing based on balanced class accuracy. Future models could prioritize authentic audio accuracy in model training and hyperparameter tuning if higher authentic accuracy is preferred.

It is important to note that the text the model was tested on had a distinct effect on the performance of the model. In authentic audio samples, the model performed better on known text for both Model-Trained and Model-Naïve participants. Conversely, in Model-Naïve clones, performance improved when the model was tested on a new paragraph. This effect was evident in both pregenerated TTS and Model-Naïve Participant clones for the Model-Trained generators. This may indicate a tendency for the model to slightly overfit to the paragraphs on which it was trained. When exposed to new participants, its performance declines. That being said, the model accuracy for authentic audio from Model-Naïve participants was 0.73. This is within half an SD of the cross-validated authentic audio accuracy (0.75, SD 0.09), further supporting the use of speech pause metrics for robust model prediction.
Incorporating features associated with real, biological processes (such as breathing, thinking, and swallowing) into a deepfake prediction algorithm is likely to enhance its reliability and longevity in the face of ongoing advancements in deepfake technologies. Instead of solely relying on a model trained on the current state of deepfake generation, which may struggle to maintain accuracy as technology evolves, the inclusion of biological features offers valuable insights that enable the model to adapt and effectively detect inauthentic voices. This approach enhances the model’s resilience against evolving deepfake techniques.

Comparison to Prior Work

High-performance current models are typically trained on spectral or deep-learned audio features obtained from the current state of deepfake generation. This permits for an extremely high accuracy in voice clones in a similar domain to the training data but new advancements and subtle changes in these obscure features could soon make these prediction models obsolete. Indeed, when a high-accuracy prediction model was tested on new, out-of-domain voice clones in a recent study, the prediction accuracy was abysmal (AUC is approximately 25%) [10]. We aimed to evaluate the use of perceptual features in current and future model implementations by testing model performance on a completely new generator. Overall, our model performance on a new generator was a success, and the average accuracy of classification of the new generator was 0.87. This generator provided no audio files for model training, and as such, we can conclude that this technique may be extended to out-of-domain cloning processes.

Limitations

This research identified certain limitations in the audio quality variation, linguistic diversity, and deepfake generators used in our study. First, since we created a new cloned audio data set, we only had a small amount of data to train and test the prediction model, and the exclusively English-focused experiments did not account for the potential impact of diverse accents or languages on our results. Small data sets may lead to model overfitting, which we attempted to mitigate using a comprehensive model testing methodology. Further exploration in this domain with a larger and more diverse data set encompassing various accents and languages is warranted, as it has the potential to strengthen the robustness of our conclusions and provide a more comprehensive understanding of model performance across linguistic variations.

Second, although the pause rate biomarker enhanced prediction accuracy, it introduced the time requirement of sufficiently long audio samples to accurately calculate pause rate data. An older data set that has been widely used for testing and training previous detection tools consisted of samples shorter than 5 seconds, rendering them incompatible with our model [19]. We prioritize the analysis of longer samples due to their higher potential for misuse in the context of misinformation or impersonation scams. Therefore, our detection tool was optimized for modern voice cloning generators and prioritized longer audio outputs over compatibility with previous deepfake data sets.

Third, another limitation concerns the variation of deepfake generation methods. Our study featured 3 distinct tools to introduce variability in deepfake audio samples. Nevertheless, numerous other models exist and possess subtle distinctions that were not covered in our investigation. While we anticipate that the incorporation of vocal biomarkers will enable accurate predictions regardless of the generation method, we did not test deepfakes produced by alternative tools. This decision stemmed from the recognition that there are numerous methods with slight variations in cloned audio samples, compelling us to focus on some of the most prominent and accessible tools.

Future Directions

In this study, we aim to show that speech pause metrics may contribute to robust deepfake detection models, and that trained models using these features perform well on out-of-domain data such as new audio deepfake generators or audio samples from new individuals. Further research should perform an ablation study to compare spectral features and pause pattern features, specifically focusing on testing on unknown data.

Conclusions

In conclusion, the integration of vocal biomarkers into machine learning models shows promise in distinguishing between authentic voice recordings and cloned samples. Given the escalating prevalence of unethical deepfake applications involving impersonation, fraud, and the dissemination of misinformation, establishing a reliable method for verifying source authenticity is crucial. Biological processes and vocal biomarkers offer a potential avenue for enhancing detection methodologies, suggesting a possible means to mitigate the risk of detection tools being rapidly outpaced by advancing deepfake generation technologies.

Acknowledgments

The authors would like to thank Klick Inc for their support in this research. The authors also thank Anirudh Thommandram for his consultation and insight on the study methodology. The authors used the generative AI tool ChatGPT by OpenAI [20] for general-purpose grammatical editing. All generated text was further reviewed and revised by the study group. No results or conclusions were impacted. The study was internally funded by Klick Inc.

Data Availability

The data sets generated or analyzed during this study are not publicly available due terms in the ethics approval and informed consent. The code used for analysis is available from the corresponding author on reasonable request.
Authors’ Contributions

NVK was responsible for conceptualization, methodology, software, investigation, data curation, and writing of the original draft and its review and editing. JK contributed to methodology, software, validation, formal analysis, investigation, and writing of the original draft and its review and editing. Additionally, JK was involved in visualization. JO participated in the methodology, investigation, and writing of the original draft and its review and editing. JO also contributed to visualization. YF played a role in the conceptualization, methodology, investigation, and writing of the original draft and its review and editing. YF also provided supervision.

Conflicts of Interest

None declared.

Multimedia Appendix 1
Speech paragraphs.

[DOCX File , 14 KB - biomedeng_v91e56245_app1.docx]

Multimedia Appendix 2
Hyperparameter tuning.

[DOCX File , 14 KB - biomedeng_v91e56245_app2.docx]

References


**Abbreviations**

ADA: AdaBoost  
AUC: area under the receiver operator curve  
DT: decision tree  
LR: logistic regression  
RF: random forest  
SVM: support vector machine  
TTS: text-to-speech  
VAD: voice activity detector

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Validation of a Novel Noninvasive Technology to Estimate Blood Oxygen Saturation Using Green Light: Observational Study

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Abstract

Background: Pulse oximeters work within the red-infrared wavelengths. Therefore, these oximeters produce erratic results in dark-skinned subjects and in subjects with cold extremities. Pulse oximetry is routinely performed in patients with fever; however, an elevation in body temperature decreases the affinity of hemoglobin for oxygen, causing a drop in oxygen saturation or oxyhemoglobin concentrations.

Objective: We aimed to determine whether our new investigational device, the Shani device or SH1 (US Patent 11191460), detects a drop in oxygen saturation or a decrease in oxyhemoglobin concentrations.

Methods: An observational study (phase 1) was performed in two separate groups to validate measurements of hemoglobin and oxygen concentrations, including 39 participants recruited among current university students and staff aged 20-40 years. All volunteers completed baseline readings using the SH1 device and the commercially available Food and Drug Administration–approved pulse oximeter Masimo. SH1 uses two light-emitting diodes in which the emitted wavelengths match with absorption peaks of oxyhemoglobin (hemoglobin combined with oxygen) and deoxyhemoglobin (hemoglobin without oxygen or reduced hemoglobin). Total hemoglobin was calculated as the sum of oxyhemoglobin and deoxyhemoglobin. Subsequently, 16 subjects completed the “heat jacket study” and the others completed the “blood donation study.” Masimo was consistently used on the finger for comparison. The melanin level was accounted for using the von Luschan skin color scale (VLS) and a specifically designed algorithm. We here focus on the results of the heat jacket study, in which the subject wore a double-layered heated jacket and pair of trousers including a network of polythene tubules along with an inlet and outlet. Warm water was circulated to increase the body temperature by 0.5-0.8 °C above the baseline body temperature. We expected a slight drop in oxyhemoglobin concentrations in the heating phase at the tissue level.

Results: The mean age of the participants was 24.1 (SD 0.8) years. The skin tone varied from 12 to 36 on the VLS, representing a uniform distribution with one-third of the participants having fair skin, brown skin, and dark skin, respectively. Using a specific algorithm and software, the reflection ratio for oxyhemoglobin was displayed on the screen of the device along with direct hemoglobin values. The SH1 device picked up more minor changes in oxyhemoglobin levels after a change in body temperature compared to the pulse oximeter, with a maximum drop in oxyhemoglobin concentration detected of 6.5% and 2.54%, respectively.

Conclusions: Our new investigational device SH1 measures oxygen saturation at the tissue level by reflectance spectroscopy using green wavelengths. This device fared well regardless of skin color. This device can thus eliminate racial disparity in these key biomarker assessments. Moreover, since the light is shone on the wrist, SH1 can be readily miniaturized into a wearable device.

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(page number not for citation purposes)
Introduction

Pulse oximetry is routinely performed in all patients with elevated body temperature. However, high blood temperature decreases the affinity of oxygen for hemoglobin (Hb) [1], and an elevation in temperature by approximately 1 °C decreases arterial oxygen saturation (sO$_2$) by only 0.5% [2]. Therefore, the decrease is very minimal and is not clinically significant in subjects with a baseline sO$_2$ level within the normal range [3].

Blood sO$_2$ measured by pulse oximetry is currently used to monitor tissue hypoxia. This method uses red and infrared wavelengths in the light spectrum and there is no correction for the level of skin melanin. Consequently, the estimated values are often inaccurate in people with darker skin tones due to the overlapping absorption spectra of melanin [4,5]. Administrative and health authorities have also recognized this issue; however, a solution has not yet been put forward [6].

To overcome this limitation, we have developed a novel technology to estimate Hb and tissue oxygenation. The scientific basis, details, and underlying technology of the device are published elsewhere [7]. In brief, the Shani device (SH1) measures reflectance of light from the skin by a pair of light-emitting diodes (LEDs), displayed as the reflectance ratio from LED1 (E1) and LED2 (E2) and as the sum of the ratios (E; E1+E2). The method of measurement is described in our previous report [7]. The light is shone on the wrist and only the reflected light is picked up by the sensors. This analog signal is then converted to a digital format by the processor, which can be analyzed, displayed, and stored in digital form (US Patent 11191460). In this device and associated technology, the reflectance ratio varies inversely with the concentration of Hb or Hb combined with oxygen (OxyHb) [8].

With an increase in body temperature, a slight drop in OxyHb is expected. Here, we focus on the results of the phase 1 study (heat jacket study) to validate the device in healthy human volunteers. The body temperature of the participants was increased by circulating warm water in a double-layered heat jacket; therefore, a slight drop in the OxyHb concentration at the tissue level was expected during the heating phase. The aim of this study was to determine whether our new device can detect the drop in sO$_2$ or a decrease in tissue OxyHb concentrations.

Methods

Investigational Device

The SH1 device was validated in tissue phantom experiments. Here, we are presenting the results of the phase 1 heat jacket study in healthy human volunteers. Details of preclinical studies and the results of these experiments have been recently published elsewhere [9]. In the preclinical experiments, we used synthetic melanin as an epilayer mimicking melanin and we used horse blood in the lower layer corresponding to the dermis of the skin. As a part of preclinical studies, we performed an absorption scan of synthetic melanin (Sigma-Aldrich, Instrument-Infinite 200 and BME089922 software system). At a melanin concentration of 1.5 mg/ml, the absorption coefficient of melanin (ie, the absorption of light per unit length) was determined to be 4.01/ml. Even with this high melanin concentration in the epilayer, our device could detect changes in OxyHb levels in the lower dermal layer. Dark-skinned subjects showed an absorption coefficient of melanin at 550 nm, corresponding to 2.5/ml. Therefore, in this preclinical study, we tested melanin levels that are higher (darker) than those measured for dark-skinned subjects. The relationship of melanin concentration with the absorption coefficient of melanin at a wavelength of 550 nm was determined in the previous study [9]. This new investigational device (SH1) measures Hb by shining light from LED1 and LED2 sequentially to obtain measures of both OxyHb and reduced Hb (ie, DeoxyHb). This is termed the “hemoglobin mode” of the operation. The reflectance ratio for LED1 is referred to as E1 and that for LED2 is referred to as E2. The reflectance ratio E1 corresponds to OxyHb, which is inversely related to the oxygen content (Figure 1). The device could detect changes in oxygen concentration in the blood, even in the presence of high melanin in the epilayer, thereby mimicking the detection of hypoxia in dark-skinned subjects.
**Eligibility and Recruitment**

As per the institutional policy, only current students and members of the staff of University of Texas at Arlington were recruited for this validation study. External candidates, including past students, were not allowed to participate. The age restriction for participation was 20-40 years. For recruitment, flyers were sent by email and posters were displayed in designated locations, including the lobby, near elevators, the cafeteria, and common rooms.

The phase 1 study was performed with two separate cohorts independently for validation of measurements of Hb and oxygen concentrations. There were 39 participants in the two studies; there was only one staff member and the remaining participants were students. Both studies included baseline measurements at visit 1. All 39 volunteers completed baseline readings using our new investigational SH1 device. Subsequently, 16 participants completed the heat jacket study and the others were included in the blood donation study. The commercially available pulse oximeter Masimo was used on the finger for comparison. Skin melanin was accounted for using the von Luschan skin color scale (VLS) and a specifically designed algorithm. Skin tone measurements were performed by two observers independently and the mean value was noted and rounded to the nearest integer. We here focus only on the results of the heat jacket study.

**Ethical Considerations**

The heat jacket study was approved by the institutional review board (IRB) at The University of Texas at Arlington (STU-2021-0150; approval date February 23, 2021). Written informed consent was obtained from each participant. Copies of consent forms are maintained by IRB authorities. The privacy and confidentiality of the participants were respected and data are stored in a coded, anonymous format. Each participant received monetary compensation as per the stipulated rules and regulations laid out by the IRB of The University of Texas at Arlington.

**Study Design**

The heat jacket study included 16 volunteers and was performed under the IRB-approved protocol. At visit 1, baseline measurements of Hb and oxygen concentrations were taken using our novel SH1 device and the commercial pulse oximeter Masimo. At visit 2, after obtaining appropriate consent, the baseline demographic information was obtained. The participant was then asked to wear a double-layered heat jacket and a pair of trousers, which comprise a network of polythene tubules and an inlet and outlet. Subsequently, the participant swallowed a telemetry pill with some water; this is a small pill-shaped electronic object that is used to sense temperature. After a few minutes, the pill reaches the stomach, measures internal body temperature, and emits a signal. A sensor attached to the jacket receives these signals, which are then relayed to a monitor via a cable. Baseline readings were taken with the participant lying down. Warm water was circulated in the jacket and trousers to increase the body temperature by 0.5-0.8°C above the baseline body temperature as measured with the telemetry pill. This increase in body temperature simulates clinically relevant fever settings. The heating phase lasted for 40 minutes, followed by cooling for the next 20 minutes. Cooling was achieved by circulating cold water through the jacket. Baseline and serial readings were taken with the new investigational SH1 device at 10-minute intervals. The measurements taken with the Masimo Pronto pulse oximeter were used for comparison. A total of 7 sets of observations were obtained for each participant over a period of 60 minutes. Readings in the heating phase and cooling phase were rescaled for each participant with baseline measurements taken as 100%. The percentage drop in sO₂ or the difference between maximum and minimum readings by the SH1 and Masimo devices was plotted for each participant.
**Results**

The mean age of the study population was 24.1 (SD 0.8) years. The skin tone varied from 12 to 36 on the VLS with a uniform distribution: one-third of the participants had fair skin, brown skin, and dark skin, respectively. Using a specific algorithm accounting for melanin, as determined from the VLS, and associated software, the reflection ratio for OxyHb is displayed on the screen along with direct Hb values [11]. We had baseline readings for all 39 participants (Figure 2) with our SH1 device and the commercial pulse oximeter Masimo. As seen in Figure 2, baseline oxygen concentrations of all 39 participants as measured by SH1 device readings fell within a range similar to those measured by the pulse oximeter.

In the heat jacket study, we expected a slight drop in OxyHb concentrations in the heating phase at the tissue level. Our hypothesis was confirmed after analyzing the data for the reflection ratios E1, E2, and E (E1+E2), followed by computation of OxyHb and total Hb values. Figure 3 shows the distribution of the skin tone (according to the VLS) of the participants compared to the percentage drop in oxygen concentrations in the 16 subjects participating in the heat jacket study. Our device could pick up more minor changes in OxyHb levels after a change in body temperature than possible with the pulse oximeter. The maximum drop in OxyHb concentrations picked up by our device was 6.5% compared to a drop of only 2.54% sensed by the pulse oximeter. The average change in OxyHb measured by our device was 2.98%, whereas that of the pulse oximeter was 1.33%, with a median of 3% and 1%, respectively.

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**Figure 2.** Skin tone versus oxygen saturation measured by the Shani device and Masimo pulse oximeter at baseline in all 39 participants. VLS: von Luschan skin coloration scale.
Discussion

Principal Findings

The baseline readings taken by the SH1 device (Figure 2) while the participants were lying in supine position showed some interesting features, including different readings for subjects with the same skin tone (e.g., subjects with skin tones 15, 16, 22, and 26 on the VLS) and similar readings for subjects with different skin tones (e.g., subjects with skin tones 15, 16, and 26 on the VLS). Therefore, the baseline measurement of tissue oxygen levels (i.e., OxyHb levels) as determined by the SH1 device is independent of skin tone and likely depends on tissue metabolism and the basal metabolic rate.

In this heat jacket study, all participants showed a drop in OxyHb levels during the heating phase, followed by a rise in the cooling phase. In some cases, the rise was slightly higher than baseline levels.

We compared the difference in maximum and minimum readings obtained by our SH1 device and the commercial pulse oximeter Masimo. Figure 3 shows the distribution of the difference across different skin tones. This drop in oxygen saturation was magnified by the SH1 device compared to the readings obtained with the pulse oximeter. The difference between maximum and minimum readings by the SH1 device was 6.5%, whereas that for Masimo was only 2.54%.

As stated earlier, the decrease in arterial oxygen levels with fever is very small, even in patients in the intensive care unit, and this drop in sO\textsubscript{2} is not clinically significant in patients with baseline sO\textsubscript{2} within the normal range. Nonetheless, this minor change is picked up well by our device and is not detected by the pulse oximeter. The commercial pulse oximeter Masimo Pronto measures arterial sO\textsubscript{2} by photoplethysmography and red-infrared wavelengths [12], whereas the novel SH1 device measures sO\textsubscript{2} at the tissue level by reflectance spectroscopy and using two green wavelengths. These are two important fundamental differences between these technologies. Table 1 compares the properties of the SH1 device and pulse oximeter.
Table 1. Comparison of the properties of the new investigational device (Shani device) and a conventional pulse oximeter.

<table>
<thead>
<tr>
<th>Properties</th>
<th>Pulse oximeter</th>
<th>Shani device</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiation used</td>
<td>Red and infrared wavelengths</td>
<td>Green visible light</td>
</tr>
<tr>
<td>Method</td>
<td>Transmittance of light</td>
<td>Reflectance of light</td>
</tr>
<tr>
<td>Any solution for melanin interference?</td>
<td>None so far</td>
<td>Yes; accounted for using a special algorithm</td>
</tr>
<tr>
<td>Accuracy of (O_2%) in dark-skinned subjects?</td>
<td>Doubtful; erratic results are obtained with darker skin tones</td>
<td>High accuracy in all subjects, regardless of skin color</td>
</tr>
<tr>
<td>Diagnostic ability (eg, hypotension, shock)</td>
<td>Poor accuracy</td>
<td>High accuracy</td>
</tr>
<tr>
<td>Continuous monitoring possible</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Site of testing</td>
<td>Fingertip</td>
<td>Back of wrist</td>
</tr>
<tr>
<td>Can be transitioned into a wearable?</td>
<td>Unclear</td>
<td>Certainly</td>
</tr>
<tr>
<td>Data storage and transfer</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Battery operated</td>
<td>Yes</td>
<td>Yes</td>
</tr>
</tbody>
</table>

Working of the Device

The emitted wavelengths of the two LEDs matched with the absorption peaks of OxyHb and DeoxyHb (ie, reduced Hb) between 520 nm and 580 nm. As mentioned above, in our new device, the input from LED1 and LED2 of the probe is received as an analog signal. This signal is then converted to digital format by the processor, which can be analyzed, displayed, and stored in digital form. In tissue phantom experiments, we bubbled air and measured the increase in oxygen concentrations in horse blood using an oxygen sensor [9]. The reflection ratio \(E_1\) received from LED1 correlates inversely with OxyHb, \(E_2\) from LED2 correlates with DeoxyHb, and \(E\) (sum of \(E_1\) and \(E_2\)) correlates with total Hb. After analyzing data from the tissue phantom experiments, we derived another new parameter termed the “OxyHb index” as a measure of OxyHb concentration. In our earlier tissue phantom experiments [9], we found that this OxyHb index varies directly with the oxygen concentration in the blood (Figure 4). This means that low oxygen concentrations are reflected as a low OxyHb index and vice versa.

Tissue oxygenation parameters include the concentrations of Hb and OxyHb in the tissue [13]. Tissue \(sO_2\) monitoring is a relatively new technology, and a drop in tissue \(sO_2\) is an early warning sign of peripheral hypoperfusion and the onset of tissue hypoxia [14]. All oximeters currently available using red and infrared wavelengths to target arterial blood flow, which measures \(sO_2\) in the conducting vascular/arterial system. Our device targets the capillary-venous network and measures tissue oxygenation. Sepsis and shock result in disturbances in microcirculatory perfusion and a change in tissue oxygen utilization that may not be reflected in arterial \(sO_2\) levels. According to many authorities, tissue oxygenation is a better marker of the underlying pathological processes as well as responsiveness to some treatments [15,16]. Tissue oxygen levels are more important, because the arterial \(O_2\) content and an adequate bulk transport of oxygen by the cardiovascular system may not guarantee delivery of oxygen to the critical tissues of the body [17]. Additionally, tissue Hb \(sO_2\) has been determined to be a better predictor of the multiorgan failure outcome [18]. Near-infrared spectroscopy also has well-known limitations [19]. To overcome these limitations, our new device uses green light (520-570 nm) for the estimation of Hb and OxyHb, while accounting for the impact of skin color on the measurements. A summary presentation displays relevant information about this technology in Multimedia Appendices 1 and 2. The video demonstrates the working of this new device.

Using green wavelengths for measurements of Hb and oxygen concentrations with a special algorithm to account for melanin is a novel concept and our efforts have already been appreciated by experts in the field [20,21]. In this study, we have used the VLS for the measurements of skin tone. This is an interval scale with measurements such as 18, 23, and 36. However, we are developing a technology to quantify melanin in the skin more precisely in a noninvasive manner, providing measurements on a continuous scale (eg, 15.25, 17.50, and 30.75). This method is patented and under development.
Strengths and Limitations
The main limitation of this study is the small sample size. Although our device fared better than the pulse oximeter, larger studies are needed in patients with diverse skin tones and a variable degree of tissue hypoxia for further clinical development of the device.

The main strength of this study is that the SH1 device offers an early warning system. A drop in tissue sO\(_2\) is an early warning sign of peripheral hypoperfusion and the onset of tissue hypoxia. Our device can sense this before traditional oximeters can raise the alarm, thereby demonstrating the potential for saving lives.

Our device can also be used in diverse settings, from home monitoring to intensive care, making critical data readily available. Light is shone on the wrist of the individual and is then measured. Therefore, in the future, this device can be miniaturized for wearable technology. The device can be operated in either hemoglobin or oximetry mode. In hemoglobin mode, the device can be used for noninvasive measurements of Hb, whereas in oximetry mode, the device can be used for the continuous monitoring of sO\(_2\).

Racial equity is another important advantage of our device. Unlike most pulse oximeters on the market, our device works for all skin tones, eliminating bias and improving care for people of color.

The specific technology underlying the design of our device offers specific advantages. First, the use of green light (520-570 nm) can obtain information from the microcirculation, revealing tissue oxygen levels invisible to red and infrared wavelengths of light. Second, our unique algorithm accounts for variations in melanin, ensuring accurate readings. These advantages can consequently lead to improved outcomes, as early detection and tailored therapies lead to better health for all, especially marginalized groups. Moreover, the ability to obtain a faster diagnosis and intervention can save both resources and lives. Finally, the device offers global reach as it is affordable and adaptable, thereby demonstrating potential to improve health care in resource-limited settings.

Conclusion
The changes in sO\(_2\) at the tissue level in normoxemic subjects are very minor or minimal. Current pulse oximeters, limited to red and infrared wavelengths, only capture the “big picture” of arterial blood flow by measuring oxygen saturation in the conducting arterial system while missing critical changes in the microcirculation, where sepsis and shock wreak havoc, before the arterial oxygen dips.

Our device is a game-changer for measuring tissue oxygenation by shining green light and offering a more sensitive marker of these hidden dangers. Our device was validated to accurately measure tissue oxygen levels and could pick up very minor changes after a change in body temperature in the heat jacket study, demonstrating improved performance compared to the commercial pulse oximeter. Tissue oxygenation parameters include the concentrations of Hb and OxyHb in the tissue. Our device worked better and appeared to be more sensitive than the pulse oximeter even for subjects with light skin or skin tone (eg, VLS 12-18). Since the sample size of this study was small, additional studies with large sample size, a diverse population, and varied degree of hypoxia are required for further confirmation.

Acknowledgments
We thank Dr Michael Nelson, Associate Professor, Department of Kinesiology, University of Texas at Arlington for allowing the use of the Kinesiology Laboratory for the clinical study. The project was funded by Shani Biotechnologies (to GA).
Data Availability
Since this is a patented technology, the data cannot be widely shared. However, the data are available with the institutional review board of the University of Texas at Arlington and can be accessed through Shani Biotechnologies LLC after completing nondisclosure agreement or confidential disclosure agreement formalities.

Authors' Contributions
SGG and GA contributed to the conceptualization, formal analysis, and methodology. VD and GA acquired the funding. GA contributed to project administration and supervision. All authors wrote the original draft and reviewed and edited the manuscript.

Conflicts of Interest
None declared.

Multimedia Appendix 1
Summary Presentation
[PPTX File, 5624 KB - biomedeng_v9i1e46974_app1.pptx]

Multimedia Appendix 2
Shani Biotech Demo(10mb)
[MP4 File (MP4 Video), 9402 KB - biomedeng_v9i1e46974_app2.mp4]

References
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Abbreviations

- DeoxyHb: deoxyhemoglobin
- Hb: hemoglobin
- IRB: institutional review board
- LED: light-emitting diode
- OxyHb: oxyhemoglobin
- SH1: Shani device
- sO2: oxygen saturation
- VLS: von Luschan skin color scale

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Preliminary Assessment of an Ambulatory Device Dedicated to Upper Airway Muscle Training in Patients With Sleep Apnea: Proof-of-Concept Study

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Abstract

Background: Obstructive sleep apnea/hypopnea syndrome (OSAHS) is a prevalent condition affecting a substantial portion of the global population, with its prevalence increasing over the past 2 decades. OSAHS is characterized by recurrent upper airway (UA) closure during sleep, leading to significant impacts on quality of life and heightened cardiovascular and metabolic morbidity. Despite continuous positive airway pressure (CPAP) being the gold standard treatment, patient adherence remains suboptimal due to various factors, such as discomfort, side effects, and treatment unacceptability.

Objective: Considering the challenges associated with CPAP adherence, an alternative approach targeting the UA muscles through myofunctional therapy was explored. This noninvasive intervention involves exercises of the lips, tongue, or both to improve oropharyngeal functions and mitigate the severity of OSAHS. With the goal of developing a portable device for home-based myofunctional therapy with continuous monitoring of exercise performance and adherence, the primary outcome of this study was the degree of completion and adherence to a 4-week training session.

Methods: This proof-of-concept study focused on a portable device that was designed to facilitate tongue and lip myofunctional therapy and enable precise monitoring of exercise performance and adherence. A clinical study was conducted to assess the effectiveness of this program in improving sleep-disordered breathing. Participants were instructed to perform tongue protrusion, lip pressure, and controlled breathing as part of various tasks 6 times a week for 4 weeks, with each session lasting approximately 35 minutes.

Results: Ten participants were enrolled in the study (n=8 male; mean age 48, SD 22 years; mean BMI 29.3, SD 3.5 kg/m²; mean apnea-hypopnea index [AHI] 20.7, SD 17.8/hour). Among the 8 participants who completed the 4-week program, the overall compliance rate was 91% (175/192 sessions). For the tongue exercise, the success rate increased from 66% (211/320 exercises; SD 18%) on the first day to 85% (272/320 exercises; SD 17%) on the last day (P=.05). AHI did not change significantly after completion of training but a noteworthy correlation between successful lip exercise improvement and AHI reduction in the supine position was observed (R=−.76; P=.03). These findings demonstrate the potential of the device for accurately monitoring participants’ performance in lip and tongue pressure exercises during myofunctional therapy. The diversity of the training program (it mixed exercises mixed training games), its ability to provide direct feedback for each exercise to the participants, and the easy measurement of treatment adherence are major strengths of our training program.

Conclusions: The study’s portable device for home-based myofunctional therapy shows promise as a noninvasive alternative for reducing the severity of OSAHS, with a notable correlation between successful lip exercise improvement and AHI reduction, warranting further development and investigation.
Introduction

Obstructive sleep apnea/hypopnea syndrome (OSAHS) is a common condition that affects a large portion of the world population [1]. It is estimated that mild to severe OSAHS affects 24% of men and 9% of women in North America [2], with an increase in prevalence over the last 2 decades [3]. OSAHS originates from repetitive closure of the upper airway (UA). The negative impacts of OSAHS include a deterioration of quality of life [4] and an increase in cardiovascular and metabolic morbidity [5-10]. Currently, the gold standard for treatment of this condition is continuous positive airway pressure (CPAP) [5] machines, which provide constant pressure to the sleeping patient via an oral or nasal mask. While this method has proven to be effective in reducing the adverse effects of OSAHS, it has been reported that from 46% to 83% of patients do not adhere to the treatment [11]. The causes of this low adherence rate may include treatment unacceptability, general discomfort, side effects (mask leaks, pressure intolerance, skin irritation, mouth dryness), bed partner intolerance, or a combination of these causes [12].

Alternatively, therapies targeting the UA muscles have been developed to decrease the disease severity [13-15]. Myofunctional therapy is a noninvasive approach in which patients are tasked with exercises of the lips, tongue, or both to target oropharyngeal functions [16]. It has been observed that myofunctional therapy may decrease the apnea-hypopnea index (AHI) by 50% in adults and by 62% in children [16]. For myofunctional therapy to be effective, the patient must perform the exercises daily. However, monitoring the quality and frequency of the exercises is pivotal to supporting implantation of such treatment and may be challenging outside of the laboratory setting. Therefore, there is a need for developing a home reeducation setup where patients can perform daily exercises with continuous monitoring of program adherence and exercise performance.

We developed a portable device that allows completion of tongue and lip myofunctional therapy while providing precise performance monitoring of performance and adherence to exercise. The aims of this clinical study were to evaluate task performance and treatment adherence to a 4-week training session and its efficacy in improving sleep-disordered breathing.

Methods

Study Design

Ten patients with untreated mild or moderate OSAHS who were referred to our sleep clinic volunteered to participate in this study. These patients were men and postmenopausal women aged ≤65 years who had a BMI ≤30 kg/m² and regular sleep habits free of sleep debt (caused by, eg, insomnia or sleep deprivation). Their initial OSAHS diagnosis and severity were established by conventional sleep studies (level 1 or 3) performed at our local sleep clinic. Consecutive patients fulfilling the entry criteria were offered enrollment, and recruitment was completed within 6 months. A polysomnographic study (level 2, Embia Titanium; Natus) and the Epworth Sleepiness Scale (ESS) were completed just before and after a 4-week training program. A registered sleep technician who was blind to the protocol performed polysomnography scoring according to standard American Academy of Sleep Medicine criteria [17]. The participants were asked to perform the full training (35 minutes) 6 days a week for 4 weeks. The first session was completed in our research laboratory and the remainder were done at the participant’s home. A follow-up was completed by phone on the first and third home training days during the first week and once a week thereafter.

Ethical Considerations

The Ethics Review Board of Institut Universitaire de Cardiologie et de Pneumologie de Québec approved the protocol (2020-3246), which conformed to the guidelines set forth by the Declaration of Helsinki, and written informed consent was obtained from all participants.

Module Overview

A briefcase-sized module that can precisely monitor lip and tongue pressure with a custom-made mouthpiece is presented in this paper. The module presented in Figure 1 is composed of 3 main parts: the mouthpiece, the interface electronics, and the user interface. The mouthpiece design is based on a 3D scan of the patient’s teeth and has 2 separate embedded internal cavities to record lip and tongue pressure. The mouthpiece is made out of silicone cast in 3D-printed sugar molds. The pressure developed by contraction of the lips or tongue is read from inside the mouthpiece with transducers and transferred to a computing unit. The module includes a touch screen with an intuitive interface for user interaction. The software combines precise pressure measurements with user calibration and engaging games to maximize therapy adherence. The training presented in the software is based on the tongue-protrusion task program presented by Svensson et al [18], in which the participants are asked to exert a force with their tongue on a force transducer and maintain a certain level of force for 1.5 seconds before releasing. The module is packaged in a customized briefcase for a robust and easy-to-transport solution.
Mouthpiece Design

To maximize adherence to the treatment, a custom mouthpiece was made for each participant. A 3D scan was made of a participant’s teeth by a dentist, and the dental print was digitally removed from a mouthpiece template using Meshmixer (version 3.5.474; Autodesk). Having a custom cutout allowed the mouthpiece to clamp naturally onto the patient’s teeth and gums, as presented in Figure 2A. The mouthpiece design includes 2 distinct internal cavities acting as pressure chambers, one for the lips and one for the tongue, as presented in Figures 2B and 2C. The lip cavity is located in the front part of the mouthpiece where the lips naturally rest. The cavity has a thin bottom and top wall where the pressure from the lips is applied. The second cavity is located in the back portion of the mouthpiece (behind the incisors) and has a thin back wall where the tip of the tongue is positioned during tongue exercises. Both cavities have tunnels connecting them to the front of the mouthpiece, where connectors can be installed to 2 distinct pressure transducers. The changes in cavity volume produced by thin wall deformation from lip or tongue movements increase the respective inner pressure.

From the digital model of the mouthpiece, a mold was created with sugar using a custom 3D printer and molding technique [19-21]. Silicon was poured into the mold, and air bubbles were removed in a vacuum chamber. Once the silicon solution solidified, the sugar was dissolved in water to free the mouthpiece.

Hardware Description

The absolute pressure in the lip and tongue cavities is linked to 2 pressure transducers (Omega PX142-002D5V) by EVA tubing and adaptors. The output voltage of the transducers is then adapted to maximize the operating range of the analog-to-digital converter (Labjack U3-LV). A computing unit (NUC7i7BNH) is used to read the values from the 2 converters. The user interacts with the software on the computing unit with a touch screen (Waveshare; this unit uses a 10.1-inch HDMI-connected LCD). All the described hardware, as well as power units, are packaged inside a briefcase, as shown in Figure 3.
Software Description

The software comprises 3 main parts: initialization, training, and games. The participant must go through a sequence alternating between these 3 parts to complete the session. Figure 4 presents a flow chart describing the process of a training session.

Zero Calibration

Once the mouthpiece has been installed, an initialization step must be performed. The participant is asked to release pressure from the tongue and lips to measure the baseline values with the display (presented in Multimedia Appendix 1, Figure S1). This step is very important since the pressure in the cavities is also influenced by temperature variation. When the participant first puts the mouthpiece on, it will gradually heat up until a steady state is reached after approximately 1 minute. The first zero calibration is followed by a 2-minute game that does not require precision in the pressure measurements. This game allows the mouthpiece temperature to reach a steady-state level. A second zero calibration takes place at the end of the game. The zero calibration step is automatically repeated before each training session to ensure that no new temperature offset affects the pressure measurements.

Maximal Pressure Calibration

After the first zero calibration and the warm-up game, the participant is asked to set a baseline for both tongue protrusion and lip contraction maximal pressure. The participant starts on a page with a vertical bar corresponding to the actual developed tongue pressure, as presented in Multimedia Appendix 1, Figure S2. The participant is asked to develop maximal tongue pressure before releasing and then press OK in 3 consecutive attempts. If the absolute deviation divided by the mean value of the maximal value is greater than 7.5%, the process is repeated; otherwise, the mean value is used as the baseline maximal tongue pressure. Subsequently, the same steps are completed for the lips.

Training

The training section consists of 3 tasks targeting, respectively, the tongue, the lips, and control of breathing. The tongue and
lip pressure tasks are based on the work of Svensson et al [18]. The display includes a white background, a green box, and an orange square, as presented in Multimedia Appendix 1, Figure S3. The orange square displays the pressure level applied to the corresponding cavity. The participant is tasked with placing the orange square into the green box. The cycle starts with the green box at the bottom for the first 8 seconds, and the participant is asked to release the pressure from the cavity. The green box then rises to a level of pressure between 3% and 5% of the maximal pressure for 3 seconds. The first 1.5 seconds allow the user to react and adjust muscle contraction to fit within the targeted pressure range. A success score corresponding to the percentage of time the participant successfully applied pressure in the given range is computed during the last 1.5 seconds. The task repeats for 10 cycles, after which a cumulative score is computed.

For the breathing task, the participant is asked to continuously apply a small lip pressure of 4% of the maximal value. Visual feedback on the lip pressure is given to the user via a vertical bar. The participant is tasked with following a breathing pattern indicated on the screen. This task includes 7 breathing cycles, with 8 seconds for inhaling and 8 seconds for exhaling. The success of this task is defined by the fraction of the total time when the user has applied sufficient pressure. The user progression during this exercise is displayed with a horizontal progress bar, as presented in Multimedia Appendix 1, Figure S4.

**Games**

The games are intended as relaxing activities between the formal metered training exercises described above. The first game is a paddle and ball game where a ball bounces off the top and bottom walls as well as paddles on the side. The user can move the paddles up in proportion to the pressure applied to the lip (blue paddle) and tongue (green paddle) cavities, respectively. The objective is to prevent the ball from hitting the edges behind the paddles. Every time the ball bounces off the paddle, a counter is incremented and the ball speed up. The counter resets once the ball hits the edge behind a paddle. The user can track the high score of the current session on the screen, as presented in Multimedia Appendix 1, Figure S5.

The second game is called the circle game. The user controls the position of a point; lip pressure controls the horizontal position and tongue pressure controls the vertical position. A yellow circle appears on the screen, and the user must combine lip and tongue pressure to place the point inside the circle. The circle turns green once the point is within its radius and must remain green for 3 seconds to succeed. Subsequently, the circle shrinks and appears at a new position. The user has 30 seconds to place the point steadily in the circle before the score resets; 7 seconds are added to the timer after each success. The current session’s high score is also displayed on-screen. The circle game display is presented in Multimedia Appendix 1, Figure S6.

In the missile game, the user must apply a small degree of tongue pressure and then release it to send a colored missile. The missile travels upward to a circle with 6 equal sections of different colors. Changing lip pressure allows the user to rotate the circle so that when the missile touches the circle, it collides with the section with the matching color. For each success, the missile velocity increases. The missile game display is presented in Multimedia Appendix 1, Figure S7.

The fourth game is the wall game, in which a ball moves horizontally toward a wall with a hole. The user must move the hole to let the ball through. The hole moves up incrementally when the user presses then releases lip pressure and moves down in the same way with tongue pressure. After each success, a new ball appears at a new height with a greater speed. The current and high scores are displayed on-screen, as presented in Multimedia Appendix 1, Figure S8.

**Results**

Once the user goes through the full training session (ie, 4 breathing tasks, 4 lip tasks, 4 tongue tasks, and 4 games), the results of each task are presented as a bar graph and are archived in a file.

**Outcomes**

The primary outcome of this study was the degree of completion of the exercises and adherence to treatment. The secondary outcomes were changes in sleep and breathing variables following training.

Compliance was defined as the number of completed sessions during the 4 weeks divided by 24 (6 sessions per week for 4 weeks). Tongue, lip, and breathing exercise success rates were defined by the percentage of time the participant successfully applied pressure in the given range. Results are presented with a 96% CI (1.96 SD). A mixed model was defined using a random intercept for the analysis of the changes in anthropometric data, the ESS score, sleep data, and the exercise success rate. One factor was associated to the before-and-after-intervention comparison. As the data were correlated, the normality assumption was verified with the Shapiro-Wilk test using residuals from the statistical model and transformed with the Cholesky metric. The graphical representation of marginal linear predictors with studentized residuals suggests the homogeneity of variances. Statistical significance was defined as a 2-tailed $P < .05$. Associations between AHI and adherence, as well as success rates, were assessed with Spearman correlations. Analyses were performed using SAS (version 9.4; SAS Institute).

**Results**

The characteristics of our study population are displayed in Table 1. This study included 2 female participants. Breathing disturbances were mostly of moderate severity, except for 1 participant who had severe sleep apnea documented during the pretraining home sleep recording. A total of 2 of the 10 recruited participants did not complete the 4-week training program due to a lack of motivation. The remainder of the participants successfully completed at least 75% (18/24) of all sessions, as presented in Table 2, with an average compliance of 91% (175/192).

For the tongue exercise, the success rate increased from 66% (211/320; SD 18%) on the first day to 85% (272/320; SD 17%) on the last day ($P = .05$). For the lip exercise, it increased from
78% (248/320; SD 18%) on the first day to 87% (278/320; SD 16%) on the last day ($P=.25$), as presented in Table 3. It is important to note that for both exercises the success score decreased in participant 2 while it improved for all other participants, except for the lip exercise for participant 7. The success rate of the breathing exercise increased from 86% (275/320; SD 24%) the first day to 96% (307/320; SD 10%) on the last day ($P=.24$).

Table 1. Anthropometric, symptom, and sleep characteristics before and after 4 weeks.

<table>
<thead>
<tr>
<th></th>
<th>Pretraining value, mean (96% CI)</th>
<th>Posttraining value, mean (96% CI)</th>
<th>$P$ value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>48 (26 to 70)</td>
<td>N/A$^a$</td>
<td>N/A</td>
</tr>
<tr>
<td>BMI (kg/m$^2$)</td>
<td>29.3 (25.8 to 32.8)</td>
<td>29.3 (26.0 to 32.6)</td>
<td>.73</td>
</tr>
<tr>
<td>ESS$^b$ score</td>
<td>11.3 (–0.1 to 22.7)</td>
<td>9.5 (–1.7 to 20.7)</td>
<td>.23</td>
</tr>
<tr>
<td>AHI$^c$ (events/h)</td>
<td>20.7 (2.9 to 38.5)</td>
<td>17.7 (–3.5 to 38.9)</td>
<td>.10</td>
</tr>
<tr>
<td>AHI supine (events/h)</td>
<td>33.4 (–3.4 to 70.2)</td>
<td>26.7 (–1.6 to 54.0)</td>
<td>.37</td>
</tr>
<tr>
<td>AHI RMI$^d$ (events/h)</td>
<td>25.5 (–6.1 to 57.1)</td>
<td>23.3 (–15.1 to 61.7)</td>
<td>.49</td>
</tr>
<tr>
<td>ODI$^e$ (events/h)</td>
<td>20.3 (–6.0 to 46.6)</td>
<td>19.2 (–9.0 to 47.4)</td>
<td>.66</td>
</tr>
<tr>
<td>Time supine (%)</td>
<td>43.7 (–19.0 to 106.4)</td>
<td>43.6 (–19.1 to 106.3)</td>
<td>.99</td>
</tr>
<tr>
<td>TST$^f$ (minutes)</td>
<td>420 (347 to 493)</td>
<td>395 (268 to 522)</td>
<td>.17</td>
</tr>
<tr>
<td>TST &lt;90% SaO$_2^g$ (%)</td>
<td>1.6 (–0.9 to 4.1)</td>
<td>1.7 (2.0 to 5.4)</td>
<td>.75</td>
</tr>
</tbody>
</table>

$^aN/A$: not applicable.

$^b$ESS: Epworth Sleepiness Scale.

$^c$AHI: apnea-hypopnea index.

$^d$RMI: respiratory mechanic instability.

$^e$ODI: oxygen desaturation index.

$^f$TST: total sleep time.

$^g$SaO$_2$: oxygen saturation of arterial blood.

Table 2. Number of training sessions completed during the 4-week training program.

<table>
<thead>
<tr>
<th>Participant</th>
<th>Week 1</th>
<th>Week 2</th>
<th>Week 3</th>
<th>Week 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (total=24)</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>2 (total=20)</td>
<td>5</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>3 (total=24)</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>4 (total=22)</td>
<td>6</td>
<td>5</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>5 (total=22)</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>6 (total=23)</td>
<td>6</td>
<td>6</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>7 (total=18)</td>
<td>4</td>
<td>3</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>8 (total=22)</td>
<td>6</td>
<td>4</td>
<td>6</td>
<td>6</td>
</tr>
</tbody>
</table>
Table 3. Success rate at baseline and after 4 weeks of training.

<table>
<thead>
<tr>
<th>Participant</th>
<th>Tongue success rate, %</th>
<th>Lip success rate, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Posttraining</td>
</tr>
<tr>
<td>1</td>
<td>34</td>
<td>82</td>
</tr>
<tr>
<td>2</td>
<td>73</td>
<td>47</td>
</tr>
<tr>
<td>3</td>
<td>61</td>
<td>99</td>
</tr>
<tr>
<td>4</td>
<td>53</td>
<td>79</td>
</tr>
<tr>
<td>5</td>
<td>62</td>
<td>90</td>
</tr>
<tr>
<td>6</td>
<td>73</td>
<td>95</td>
</tr>
<tr>
<td>7</td>
<td>92</td>
<td>95</td>
</tr>
<tr>
<td>8</td>
<td>79</td>
<td>92</td>
</tr>
</tbody>
</table>

During the study period, there was no significant change in BMI, ESS score, AHI, and other polysomnography-derived parameters, as displayed in Table 1. Table 4 presents the variation in AHI before and after the 4-week training for each participant. It illustrates that the index of each participant improved, with the exception of participants 2 and 3. A significant correlation was found between the decrease in AHI in the supine position and the change in success rate for the lip exercise ($R_s=-0.76; P=.03$).

Table 4. Effects of 1 month of training on obstructive sleep apnea/hypopnea syndrome severity.

<table>
<thead>
<tr>
<th>Participant</th>
<th>Pretraining AHI, events/h</th>
<th>Posttraining AHI, events/h</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>13.6</td>
<td>11.6</td>
</tr>
<tr>
<td>2</td>
<td>15.4</td>
<td>19.6</td>
</tr>
<tr>
<td>3</td>
<td>41.0</td>
<td>40.8</td>
</tr>
<tr>
<td>4</td>
<td>17.4</td>
<td>6.3</td>
</tr>
<tr>
<td>5</td>
<td>25.0</td>
<td>21.3</td>
</tr>
<tr>
<td>6</td>
<td>20.5</td>
<td>14.9</td>
</tr>
<tr>
<td>7</td>
<td>12.6</td>
<td>8.3</td>
</tr>
<tr>
<td>8</td>
<td>20.2</td>
<td>19.0</td>
</tr>
</tbody>
</table>

Discussion

Principal Findings

The results of this study illustrate the feasibility of performing a training task focusing on the recruitment of different UA muscles while collecting major information such as adherence to the training program and objective measurements of task completion and success.

Comparison to Prior Work

Myofunctional therapy is a relatively new treatment for sleep-disordered breathing and is based on a combination of regular exercises aiming at enhancing muscle recruitment from various oral and oropharyngeal structures [13]. Although its effect on AHI and sleep apnea–related symptoms has been shown in an increasing number of studies [16], the main challenge to its success remains the objective assessment of program adherence and exercise performance [22]. Of the 10 participants initially recruited, 8 successfully completed more than 75% (18/24) of the total number of sessions over a period of 4 weeks. Two of the initially recruited participants left the study less than 3 days after the start of the training. In comparison, rates of participant adherence reported in recent studies using myofunctional therapy with a mobile app were 75%, 90%, and 65% (15 minutes per session, 5 times/week for 3 months) [23-25]. In addition, Kim et al [26] found that in a myofunctional therapy support program with the help of exercise diaries, the reported adherence was 82%. In studies from Kim et al [26] and O’Connor et al [23-25], one could note the strong encouragement given to the patients through easy access to a health professional, encouraging text messages, or the use of a mobile app. In this study, a higher adherence rate was observed compared to the aforementioned studies. One potential explanation for this improvement is that the exercises were designed to induce motivation with games and visual feedback. Recently, a mobile app has been developed for this purpose that only requires a smartphone [24]. It first teaches the patient how to perform the exercise and then provides timely feedback on their performance. The results are saved over time, and the app promotes assiduity. It was observed that after 3 months, 75% of the patients completed the training at least 5 days a week [24] and the AHI of patients who adhered to the treatment decreased by 53.4% [23]. However, the quality tracking of the
exercises is limited by the functionality of the smartphone and requires covering the screen with cling film or hypoallergenic plastic wrap every session, since the tongue touches the screen [24]. The proposed custom mouthpiece introduced in this paper enables more accurate measurement of lip and tongue pressure, ensuring enhanced exercise quality.

**Strengths**

The multidimensional nature (exercises mixed with various training games) of the training program, its ability to immediately provide performance results for each exercise to the participants, and the measurement of treatment adherence are important strengths of our training program. As myofunctional therapy is based on an integrative approach, it is not possible to define which of the exercises may contribute most significantly to treatment success [13]. In recent studies, a combination of 9 exercises has been found to be sufficient to significantly decrease AHI [23,24]. Previous studies have focused on a single exercise and have found variable success [14,27]. Here, we used 3 exercises as a compromise between recruiting more of the muscles involved in the pathogenesis of OSAHS and keeping the workload at an acceptable level for participant motivation. Our results seem to show that a greater number of participants should have been recruited in order to see a higher impact on AHI reduction. Based on the results of our previous study [14], the training program duration was set at 4 weeks. This may have affected the results, as clear benefits in previously published myofunctional therapy studies were observed after 3 months of training. This difference may have helped to obtain a higher adherence rate while limiting the AHI reduction.

**Limitations**

There were 3 limitations of this study. The first was the success rate. This 4-week program did not significantly influence OSAHS severity to the degree that we previously found with an intensive in-lab tongue-protrusion training session that lasted 1 hour in 1 week [14]. However, it should be emphasized that although AHI did not decrease significantly, a correlation was found between the increase in success rate for the lip exercise and AHI decrease in the supine position. Therefore, one possible explanation for the modest decrease in AHI could be that the success rate of the present exercises was much higher than our previous in-lab trial (for both success rate during the first session and the rate of increase during the training period). Previously, it started from an average of 28% up to 65% for the last session. Similarly, an initial success rate of 25% was observed by Svensson et al [18] in a similar 1-week tongue-training program that was devised to increase corticomotor excitability. The high success rate in this study was likely mainly due to the adjustments that were made to the experimental set-up in order to make it ambulatory. However, in our success rate calculation, we did not take into account the results of the game sessions, which also involved a learning process and accounted for about half the duration of a training session. These games were designed mainly to boost the patients’ motivation to continue the program. Future exercise settings could be individually adjusted to adapt exercise targets to participants’ baseline UA performance, with the goal of improving the success rate over time.

The second limitation was the selection of participants. Since it is not known to what extent anatomical UA abnormalities contribute to training program efficacy, no such selection criteria were used for our study population. It could be interesting to complete further studies focusing on patients with limited anatomical abnormalities according to practical clinical scores (ie, Mallampati and velopharyngeal scores). Apart from sleep apnea severity and degree of obesity, there were no selection criteria for participant selection. However, there may have been an indirect selection criterion due to the need for the participants to complete an additional preliminary visit at a dentist’s office to perform the 3D tooth scan. This may have interfered with individual willingness to enter into the trial. Identifying participants who will remain engaged with the training program is a crucial issue for such a treatment strategy. Having a training device available for demonstration in the setting of a sleep clinic could definitely help to identify participants who are likely to follow the requirements of a training program.

The third limitation was the sample size. Yet another explanation for the lack of an AHI decrease is that our sample size was affected by the dropout of 2 participants and by the increase in AHI observed after the intervention in 1 participant. This participant was the only one with a decreasing overall success rate. This particular patient did have difficulty remaining focused throughout the training month, probably due to excessive sleepiness. As OSAHS is a multifactorial disorder [28] in which anatomical and nonanatomical factors can interact to modulate the severity of the disease, patient selection may play an important role in the success of OSAHS muscle training. Therefore, for this particular participant, treatments targeting traits other than low muscle tone or function would have been effective for decreasing AHI and related symptoms.

**Conclusions**

This study was an attempt to develop a prototype aimed at completing a simplified oral/oropharyngeal exercise program in an entertaining way in the comfort of a patient’s own home. The program gives the patient visual feedback, as well as the ability to monitor improvement. Patients were instructed to perform tongue protrusion exercises, lip pressure exercises, and controlled breathing in various playful tasks 6 times a week for 4 weeks. Session duration was about 35 minutes. While the AHI reduction was not significant, we found that the success rate for improvement in the lip exercise was correlated with AHI reduction in the supine position ($R_s=-0.76; P=.03$). These results are a first step toward the tuning of an ambulatory myofunctional therapy module able to accurately monitor the performance of participants in lip and tongue pressure exercises. This noninvasive approach may decrease the severity of OSAHS and represent an alternative to more invasive solutions, such as CPAP devices.
Acknowledgments

The authors would like to gratefully acknowledge the Fonds Alphonse-L’Espérance and the Ministère de l’Économie, de l’Innovation et de l’Énergie du Gouvernement du Québec for the financial support they provided for this research.

Data Availability

The data sets generated during and/or analyzed during this study are available from the corresponding author on reasonable request.

Authors’ Contributions

PR contributed to writing (original draft, review, and editing), software, and visualization. JR contributed to conceptualization, project administration, funding acquisition, supervision, and writing (review and editing). AB-D contributed to supervision and writing (review and editing). JL contributed to supervision and writing (review and editing). SG contributed to data curation, formal analysis, methodology, and writing (review and editing). JFM contributed to resources and writing (review and editing). FS contributed to conceptualization, project administration, funding acquisition, supervision, and writing (review and editing).

Conflicts of Interest

None declared.

Multimedia Appendix 1

Screenshots of the software.

References


Abbreviations

- AHI: apnea-hypopnea index
- CPAP: continuous positive airway pressure
- ESS: Epworth Sleepiness Scale
- OSAHS: obstructive sleep apnea/hypopnea syndrome
- UA: upper airway

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Impact of Audio Data Compression on Feature Extraction for Vocal Biomarker Detection: Validation Study

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Abstract

Background: Vocal biomarkers, derived from acoustic analysis of vocal characteristics, offer noninvasive avenues for medical screening, diagnostics, and monitoring. Previous research demonstrated the feasibility of predicting type 2 diabetes mellitus through acoustic analysis of smartphone-recorded speech. Building upon this work, this study explores the impact of audio data compression on acoustic vocal biomarker development, which is critical for broader applicability in health care.

Objective: The objective of this research is to analyze how common audio compression algorithms (MP3, M4A, and WMA) applied by 3 different conversion tools at 2 bitrates affect features crucial for vocal biomarker detection.

Methods: The impact of audio data compression on acoustic vocal biomarker development was investigated using uncompressed voice samples converted into MP3, M4A, and WMA formats at 2 bitrates (320 and 128 kbps) with MediaHuman (MH) Audio Converter, WonderShare (WS) UniConverter, and Fast Forward Moving Picture Experts Group (FFmpeg). The data set comprised recordings from 505 participants, totaling 17,298 audio files, collected using a smartphone. Participants recorded a fixed English sentence up to 6 times daily for up to 14 days. Feature extraction, including pitch, jitter, intensity, and Mel-frequency cepstral coefficients (MFCCs), was conducted using Python and Parselmouth. The Wilcoxon signed rank test and the Bonferroni correction for multiple comparisons were used for statistical analysis.

Results: In this study, 36,970 audio files were initially recorded from 505 participants, with 17,298 recordings meeting the fixed sentence criteria after screening. Differences between the audio conversion software, MH, WS, and FFmpeg, were notable, impacting compression outcomes such as constant or variable bitrates. Analysis encompassed diverse data compression formats and a wide array of voice features and MFCCs. Wilcoxon signed rank tests yielded P values, with those below the Bonferroni-corrected significance level indicating significant alterations due to compression. The results indicated feature-specific impacts of compression across formats and bitrates. MH-converted files exhibited greater resilience compared to WS-converted files. Bitrate also influenced feature stability, with 38 cases affected uniquely by a single bitrate. Notably, voice features showed greater stability than MFCCs across conversion methods.

Conclusions: Compression effects were found to be feature specific, with MH and FFmpeg showing greater resilience. Some features were consistently affected, emphasizing the importance of understanding feature resilience for diagnostic applications. Considering the implementation of vocal biomarkers in health care, finding features that remain consistent through compression for data storage or transmission purposes is valuable. Focused on specific features and formats, future research could broaden the scope to include diverse features, real-time compression algorithms, and various recording methods. This study enhances our understanding of audio compression’s influence on voice features and MFCCs, providing insights for developing applications across fields. The research underscores the significance of feature stability in working with compressed audio data, laying a foundation for informed voice data use in evolving technological landscapes.

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KEYWORDS
vocal biomarker; biomarker; biomarkers; sound; sounds; audio; compression; voice; acoustic; acoustics; audio compression; feature extraction; Python; speech; detect; detection; algorithm; algorithms

Introduction

Background
Vocal biomarkers are emerging as a promising accessible and noninvasive avenue for medical screening, diagnostics, and monitoring [1]. These biomarkers are unique characteristics or acoustic patterns of an individual’s voice that can hold valuable information about their physical and mental well-being [2]. Human voice production requires the coordination of multiple biological systems; perturbations in these systems induced by various conditions or diseases can result in alterations in the characteristics of the human voice [3]. Potential applications of vocal biomarkers are diverse, including the identification of neurological disorders, cardiovascular diseases, respiratory conditions, and mental health disorders, among others [2,4-6].

In our previous work, “Acoustic Analysis and Prediction of Type 2 Diabetes Mellitus Using Smartphone-Recorded Voice Segments” [7], smartphone-recorded speech was used to predict type 2 diabetes mellitus through a comprehensive acoustic analysis [7]. The study demonstrated the feasibility of using acoustic features from smartphone-recorded voice data to predict the presence of this disorder, highlighting the valuable diagnostic potential of vocal biomarkers in the context of a specific health condition [7]. Building upon this prior research, we aim to assess the impact of audio compression on acoustic vocal biomarker development, which is crucial for the broader applicability of this emerging field.

The development of acoustic vocal biomarkers relies on the analysis of voice data, and this process is multifaceted. One critical aspect of this analysis is feature extraction, which involves identifying and quantifying relevant acoustic features from the voice data [8]. These features may encompass a wide range of parameters such as pitch, spectral properties, prosodic patterns, and various other characteristics that carry meaningful information about the speaker’s health status [2,8]. Accurate and robust feature extraction is pivotal for the successful identification and interpretation of vocal biomarkers.

Voice data are often captured, transmitted, and stored in various digital formats that may include compression, a common practice used to reduce the size of audio files, making them more manageable and efficient for storage and transmission [9]. It is necessary to consider the potential impact of audio data compression on the overall process of vocal biomarker development as the process can have significant effects on the audio [10]. Compression algorithms are widely applied to raw, high-quality audio (typically waveform audio file format) and can be classified as lossy or lossless [11]. Lossy compression algorithms reduce file size to as low as 10% of the original size by removing mostly inaudible audio data, while lossless preserves all the original audio data and only compresses to approximately 50% [12]. Some of the most common lossy formats include MP3, M4A, and WMA [12]. These formats offer different trade-offs between file size and audio quality, and each may introduce specific artifacts and alterations to the original acoustic data.

Previous research on how data compression impacts voice signals has found that different microphones and MP3 compression bitrates on sustained vowel sounds can significantly affect feature values [10]. Research has found that various digital platforms and their audio codecs affect the voice in a way that challenges voice recognition processes specifically by narrowing the frequency band and centrally shifting frequencies at the upper and lower limits [13]. While differing microphones can introduce differences in audio data depending on specifications, smartphone microphones have been found to collect high-quality audio data suitable for acoustic analysis [14].

This exploratory research aims to investigate the effect of common audio data compression algorithms, such as MP3, AAC (compression algorithm for M4A), and WMA, on the vocal biomarker feature extraction process. Additionally, the effect of compression bitrate or encoder type will be analyzed to determine whether these factors make a difference within each format. Understanding the impact of popular data compression methods on acoustic vocal biomarker analysis is important as it can significantly affect the quality and interpretability of biomarker data [15,16]. Moreover, this knowledge can guide the development of best practices and inform the compression implementation process for the specific needs of health care applications, such as remote medical care involving telephone or video conferencing, thereby minimizing the risk of unintentional distortion of vocal biomarkers.

Objective
The objective of this research is to analyze the effect of several common audio data compression algorithms: MP3, M4A, and WMA, in 2 common bitrates, completed by 3 different conversion tools, on feature extraction from voice data for vocal biomarker detection.

Methods
Overview
In this research, acoustic features were derived from uncompressed voice samples, which were subsequently converted into MP3, M4A, and WMA formats using 3 distinct tools, namely MediaHuman (MH) Audio Converter, WonderShare (WS) UniConverter, and Fast Forward Moving Picture Experts Group (FFmpeg) across 2 different bitrates (320 and 128 kbps). MH, WS, and FFmpeg conversion tools were selected because of their accessibility as free, downloadable audio conversion software. Our goal was to explore how different audio conversion tools, formats, and 2 specific bitrates affect the data set used to develop a biomarker prediction model [7]. By focusing on these tools and bitrates, we aimed to provide insights into the potential impact of common audio compression methods on the extracted voice features. This approach allowed for a manageable analysis while paving the way for future
research to delve deeper into the nuances of audio compression effects on biomarker prediction models.

Data and Participants
This research was conducted using a data set of audio recordings that were collected from 505 participants (mean age 41.03, SD 13.29 years, 336 male participants) recruited between August 30, 2021, and June 30, 2022, for a study in India [7]. Participants were instructed to record a short English phrase up to 6 times daily using their smartphone for 14 consecutive days. As these data were originally recorded for research involving diabetes, the phrase was “Hello. How are you? What is my glucose level right now?” All audio files used in the research originated in the uncompressed waveform audio file format, 16-bit 44.1 kHz.

Participants in this study used a variety of smartphone models for data recording. While efforts were made to request recordings in quiet environments, the inherent difficulty in controlling recording conditions may have introduced variability in the recorded speech data. No preliminary tests were conducted to assess the recording quality across different smartphone models, and no preprocessing techniques were applied to address potential hardware variations in the recorded speech data. It is noteworthy that the intention of the prediction model was to be run on a smartphone; therefore, the recordings were made using smartphone uncompressed audio to align with the intended application context.

File Conversion
To explore the impact of diverse data compression methods, the original files underwent conversion using MH (version 2.2.2), WS (version 15), and FFmpeg (version 6.1.1) in Python (version 3.10.11; Python Software Foundation) on a PC. Three distinct compression algorithms—MP3, M4A, and WMA—at 2 bitrates—128 kbps and 320 kbps—were applied to simulate real-world scenarios where audio data are commonly subjected to different compression algorithms for storage and transmission purposes. The sample rate (44.1 kHz) and the channels (stereo) were kept consistent over all formats. The choice of encoders used in the research was not a primary consideration; rather, our focus was on comparing the results obtained from different compression methods. It is worth noting that the selected encoders were accessible, free, and capable of batch processing multiple files, which facilitated efficient experimentation. Despite maintaining consistency in factors such as bitrate, channels, and formats between the 3 encoders, there are features of the tools that remain hidden that could potentially cause differences in the converted files, such as the encoding mode (ie, constant or variable bitrate) or other encoding options. However, these hidden features are not a large concern because the objective of the study was to compare compressed and uncompressed data rather than comparing between compression. The incorporation of multiple encoders served the purpose of discerning whether factors beyond just bitrate and file format influenced feature values.

Feature Extraction and Comparison
We chose to use the same feature set (Multimedia Appendix 1) as in our previous research on developing a voice-to-type 2 diabetes model to maintain consistency and leverage their established effectiveness in capturing relevant biological information from voice data [7]. Acoustic features were extracted from both the original waveform audio file format files and the compressed audio formats using Python (version 3.10.11; Python Software Foundation). The voice feature extraction process leveraged Parselmouth, a Python integration of Praat speech and voice analysis software [17,18], ensuring robustness and accuracy in feature extraction. The extracted features aimed to capture pertinent acoustic characteristics of the voice data, such as pitch, jitter, and intensity, as well as Mel-frequency cepstral coefficients (MFCCs) [19], which have demonstrated efficacy in capturing subtle variations in vocal properties associated with health conditions.

Notable perceived voice qualities such as breathiness, hoarseness, and roughness, which typically present with elevated levels of shimmer and jitter, were often associated with certain pathological conditions and were therefore included in the biomarker development as well as this research [7,20]. While acoustic analysis is mainly performed using sustained phonation of vowel sounds, recent studies have demonstrated the use of shimmer and jitter measurements in identifying dysphonia even when calculated from entire sentence recordings [20]. Thus, because the data set was originally studied for the purpose of biomarker development, we chose to include the evaluations of shimmer and jitter alongside traditional vocal parameters such as pitch, intensity, and harmonic noise ratio in this analysis of how audio data compression impacts feature values.

Given the non-Gaussian distribution of feature data, assessed via the Shapiro-Wilk test, a nonparametric approach—specifically, the Wilcoxon signed rank test—was adopted for statistical analysis. This paired test aimed to evaluate the impact of each compression method on audio features by comparing the features extracted from the original uncompressed files with those obtained from each compressed format individually. In this study, the Bonferroni correction method was used to account for multiple comparisons. Given our focus on assessing the impact of each conversion method relative to the original feature values rather than comparing between different treatments, this correction was deemed appropriate. This approach allowed us to effectively manage the potential for false positives while evaluating the stability of feature values across different compression methods.

Ethical Considerations
The protocol (ID MGCTS107) received ethics approval by Saanvi Ethical Research LLP, all participants signed informed consent, and data were stored in a secure cloud database with no identifying information. Participants were compensated for their time.

Results
Data and Participants
A total of 36,970 audio files were recorded from the 505 participants who completed the study. Speech-to-text screening ensured that the audio files adhered to the fixed sentence criteria and were devoid of substantial background noise, resulting in
File Conversion

The noncustomizable differences between the audio conversion software MH Audio Converter and WS UniConverter manifested in evident variations in the converted files. Table 1 displays the differences in compression ratio and data set size, highlighting these distinctions and emphasizing the impact of software-specific characteristics on the compression outcomes such as constant or variable bitrates.

Table 1. Compression specifications for each data compression method including the final size of the data set and compression ratio.

<table>
<thead>
<tr>
<th>Format and tools</th>
<th>Bitrate (kbps)</th>
<th>Data set size (GB)</th>
<th>Compression ratio</th>
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\(^a\)FFmpeg: Fast Forward Moving Picture Experts Group.

Feature Extraction and Comparison

This research investigated the influence of diverse data compression formats on an extensive array of voice features and MFCCs. The corresponding \( P \) values for each feature are provided in the subsequent table from the results of the 756 Wilcoxon signed rank tests. \( P \) values below the level of significance, \( 6.61 \times 10^{-5} \) with the Bonferroni correction (Table S1-S3 in Multimedia Appendix 2), signify a notable difference in feature values between the original .wav format and the corresponding compressed format, indicating a significant alteration due to compression. Conversely, features with \( P \) values greater than \( 6.61 \times 10^{-5} \) (Table S1-S3 in Multimedia Appendix 2) are deemed robust, suggesting their resilience to the compression process.

Discussion

Overview

This investigation illuminated the effects of diverse audio file compression methods on a broad spectrum of voice features and MFCCs. The results revealed that the impact of data compression is feature specific and varies across different encoders, formats, and bitrates.

Principal Findings

The encoder played a substantial role in influencing voice features, with MH- and FFmpeg-converted files demonstrating greater resilience to compression compared to WS-converted files, regardless of the format. For MH, WS, and FFmpeg, there were 15, 6, and 21 features, respectively, that had at least 1 format or bitrate combination that was unaffected by the conversion. A total of 59 compressed feature comparisons showed stability for MH, 8 for WS, and 67 for FFmpeg (Table S1-S3 in Multimedia Appendix 2). The conversion bitrate also
exhibited an impact on feature stability, with some features remaining consistent for both bitrates, while others were affected uniquely at either 128 kbps or 320 kbps. A total of 38 feature comparison cases (of the total of 134) were only affected by compression for a single bitrate. Of those 38, 15 feature comparisons were only unaffected with 128 kbps, while 23 were stable for compression at only 320 kbps. MH and FFmpeg conversions had more features unaffected when conversions were done with a bitrate of 320 kbps compared to 128 kbps. Additionally, the voice features were found to be more stable than the MFCCs. The findings indicate that not all voice features respond equally to audio file compression. Certain features exhibited robustness and remained consistent despite compression, holding promise for applications involving compressed voice data storage or transmission. For instance, in our previous work on type 2 diabetes prediction from voice, features such as mean fundamental frequency/pitch (meanF0), pitch SD (stdevF0), and relative average perturbation jitter (rapJitter) remained consistent across several compression methods, including MP3 from MH at 320 kbps and FFmpeg at both 320 and 128 kbps and WMA from MH and FFmpeg at both 128 kbps and 320 kbps (Table S1-S3 in Multimedia Appendix 2) [7]. For the male prediction model, 1 of the 2 features (meanI) was significantly affected by all conversion methods. The second feature (apq11) remained stable for conversions with MH and FFmpeg to WMA format at both bitrates, MP3 at 320 kbps, and MH-converted M4A at 320 kbps. (Table S1-S3 in Multimedia Appendix 2) [7]. However, this study also identified features significantly altered by compression (Table S1-S3 in Multimedia Appendix 2), emphasizing the need to understand the stability and sensitivity of individual features for maintaining accuracy and interpretability in applications like health care diagnostics and voice recognition.

Vocal biomarkers, being a relatively new concept, are predominantly situated within the realm of research rather than practical settings where considerations for data storage and transmission are paramount. The study’s implications extend to various fields, particularly in health care, where voice data are increasingly used for disease detection and monitoring. When dealing with features significantly influenced by a specific compression algorithm, considerations should be made to preserve accuracy in applications requiring high diagnostic precision. The study suggests that certain voice features can withstand common data compression formats, enabling the use of compressed data in medical applications without compromising diagnostic accuracy, depending on the features. This is crucial in scenarios involving limited bandwidth for audio data transmission or storage constraints, where choosing an appropriate compression format while considering feature resilience becomes pivotal. Conversely, for research applications where features are being investigated, the use of uncompressed or lossless compression is essential.

Limitations and Future Directions
This study has several limitations. First, while it focused on a specific set of voice features and how they were changed based on compression formats, future research could benefit from isolating compression settings to study their individual effects rigorously. Second, controlling microphone and recording settings could enhance data consistency and reliability, as variations in these factors may introduce confounding variables. Additionally, exploring different recording sentences could provide insights into how content variability influences the impact of compression on feature extraction. Finally, a broader exploration of diverse features beyond those examined in this study, such as spectral or temporal features, could offer a more comprehensive understanding of the impact of compression on acoustic vocal biomarkers.

Conclusions
In this research, we have provided insights into the influence of audio data compression on feature values used in biomarker prediction model development. Our findings underscore the importance of considering compression effects in the design and optimization of diagnostic tools reliant on voice-based biomarkers. Through analysis and statistical comparisons, we have demonstrated the nuanced impact of compression formats, bitrates, and conversion tools on the stability and reliability of extracted feature values. By revealing these effects, our research not only advances our understanding of the complex interplay between audio data processing and biomarker extraction but also offers practical implications for health care practitioners and researchers. Moving forward, the findings pave the way for future investigations aimed at refining compression strategies, exploring alternative extraction methodologies, and ultimately enhancing the accuracy and efficacy of biomarker-based diagnostic models in clinical practice.

Acknowledgments
This research was internally funded by Klick Inc.

Data Availability
Data were commissioned by Klick Health, a private business, and are owned by Klick who does not allow sharing them.

Authors’ Contributions
Data processing and analysis was done by JO. Feature extraction method was developed by JK. All authors participated in manuscript editing.
**Conflicts of Interest**

All authors are employees of Klick Inc.

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Multimedia Appendix 1

Voice feature set.

[DOCX File, 14 KB - biomedeng_v91e56246_app1.docx]

Multimedia Appendix 2

Pairwise comparisons for feature values between compressed and uncompressed audio data.

[DOCX File, 51 KB - biomedeng_v91e56246_app2.docx]

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**References**


Abbreviations

**FFmpeg:** Fast Forward Moving Picture Experts Group

**MFCC:** Mel-frequency cepstral coefficient

**MH:** MediaHuman

**WS:** WonderShare

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A Deep Learning Framework for Predicting Patient Decannulation on Extracorporeal Membrane Oxygenation Devices: Development and Model Analysis Study

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Abstract

Background: Venovenous extracorporeal membrane oxygenation (VV-ECMO) is a therapy for patients with refractory respiratory failure. The decision to decannulate someone from extracorporeal membrane oxygenation (ECMO) often involves weaning trials and clinical intuition. To date, there are limited prognostication metrics to guide clinical decision-making to determine which patients will be successfully weaned and decannulated.

Objective: This study aims to assist clinicians with the decision to decannulate a patient from ECMO, using Continuous Evaluation of VV-ECMO Outcomes (CEVVO), a deep learning–based model for predicting success of decannulation in patients supported on VV-ECMO. The running metric may be applied daily to categorize patients into high-risk and low-risk groups. Using these data, providers may consider initiating a weaning trial based on their expertise and CEVVO.

Methods: Data were collected from 118 patients supported with VV-ECMO at the Columbia University Irving Medical Center. Using a long short-term memory–based network, CEVVO is the first model capable of integrating discrete clinical information with continuous data collected from an ECMO device. A total of 12 sets of 5-fold cross validations were conducted to assess the performance, which was measured using the area under the receiver operating characteristic curve (AUROC) and average precision (AP). To translate the predicted values into a clinically useful metric, the model results were calibrated and stratified into risk groups, ranging from 0 (high risk) to 3 (low risk). To further investigate the performance edge of CEVVO, 2 synthetic data sets were generated using Gaussian process regression. The first data set preserved the long-term dependency of the patient data set, whereas the second did not.

Results: CEVVO demonstrated consistently superior classification performance compared with contemporary models (P<.001 and P=.04 compared with the next highest AUROC and AP). Although the model’s patient-by-patient predictive power may be too low to be integrated into a clinical setting (AUROC 95% CI 0.6822-0.7055; AP 95% CI 0.8515-0.8682), the patient risk classification system displayed greater potential. When measured at 72 hours, the high-risk group had a successful decannulation rate of 58% (7/12), whereas the low-risk group had a successful decannulation rate of 92% (9/9) (P=.01). When measured at 96 hours, the high- and low-risk groups had a successful decannulation rate of 54% (6/11) and 100% (9/9), respectively (P=.01). We hypothesized that the improved performance of CEVVO was owing to its ability to efficiently capture transient temporal
patterns. Indeed, CEVVO exhibited improved performance on synthetic data with inherent temporal dependencies ($P<.001$) compared with logistic regression and a dense neural network.

**Conclusions:** The ability to interpret and integrate large data sets is paramount for creating accurate models capable of assisting clinicians in risk stratifying patients supported on VV-ECMO. Our framework may guide future incorporation of CEVVO into more comprehensive intensive care monitoring systems.

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**KEYWORDS**
extracorporeal membrane oxygenation; ECMO; venovenous; VV; machine learning; supervised learning; dynamic data; time series; clinical decision support; artificial intelligence; AI; clinical AI; health informatics

**Introduction**

**Background**

Extracorporeal life support (ECLS) is a suite of resource-intensive therapies indicated in patients with refractory respiratory failure or cardiogenic shock [1]. This intervention involves cannulation of central or peripheral arteries and veins to provide forward flow through a circuit with a mechanical pump and gas exchange device, also called a membrane oxygenator. Air is connected to the membrane oxygenator to deliver oxygen and remove carbon dioxide from the circulating blood. Established indications for venovenous extracorporeal membrane oxygenation (VV-ECMO) exist in the literature, and the use of this technology was expanded during the COVID-19 pandemic [2]. The VV-ECMO configuration is specifically used for patients experiencing severe lung injury. This setup is designed to provide oxygenation and decarboxylation support without offering the additional hemodynamic assistance found in the venoarterial configuration. VV-ECMO is considered a last resort therapy for patients with end-stage respiratory failure [3], with an overall survival rate of 60% [4].

Decannulating a patient from VV-ECMO is a clinical challenge that requires considerable training and expertise from provider teams in the intensive care unit. Clinicians assess trends in the patient’s vital signs, physical examination, response to various therapies, laboratory biochemistries, and radiographic studies. When the decision is made to proceed, decannulation is usually accomplished through a weaning trial during which VV-ECMO is gradually reduced. To date, there are limited prognostication scores that successfully predict when patients are ready to undergo a weaning trial. In this study, we present an artificial intelligence model capable of running in real time that incorporates discrete and continuous variables that clinicians may use in their assessment of patients for decannulation from VV-ECMO support.

**Related Work**

Multiple predictive scores have been developed to help clinicians prognosticate before cannulation. The 6 most common prognostication scores for adult respiratory failure supported on extracorporeal membrane oxygenation (ECMO) are ECMOnet, Predicting Death for Severe Ards on VV-ECMO, Respiratory ECMO Survival Prediction, Roch, Venovenous ecmo mortality score, and Prediction of Survival on ECMO Therapy score [5] (Table S1 in Multimedia Appendix 1 [6-11]). Although these 6 scores are commonly used, they have 2 main drawbacks. First, all input information is recorded before cannulation to ECMO because the primary intent of the models is to be used to determine which candidates were most likely to benefit from the intervention. Second, all scores use logistic regression to predict outcomes or identify significant variables. Logistic regression requires high-quality data from static variables, which limits the types of data that can be inputted. Thus, sequential or time-series data such as laboratory values and vital signs must be limited to a single time point or summarized. Furthermore, these statistical models are limited in terms of capturing nonlinear effects and interactions between variables.

To date, no studies have mitigated both issues to improve the prediction of successful decannulation in patients supported on VV-ECMO. However, some researchers have attempted to use deep learning to predict specific clinical events. Abbasi et al [12] used clinical and ECLS data to compare 2 approaches, deep learning and traditional statistical methodology, to develop a model to predict hemorrhage and thrombosis events. The deep learning model outperformed linear regression in both hemorrhage and thrombosis data sets, suggesting that more complex models may achieve better predictive power. Other authors have applied deep learning and modified logistic regression to predict survival on venoarterial-ECMO (VA-ECMO) only. Ayers et al [13] used 48 hours of laboratory values after VA-ECMO cannulation to predict survival to discharge using a deep neural network. Similarly, Loyaga et al [14] used clinical, echocardiographic, laboratory, and hemodynamic characteristics to predict 30-day mortality in patients on VA-ECMO using the elastic-net method. None of these studies used data obtained from the ECMO devices, and instead used laboratory values, clinical scores, and disease severity to train their models. These approaches leave a large amount of valuable information unused. In the hospital, clinicians adjust the parameters of ECMO support in real time according to the patient’s condition and pathophysiology. Modern devices capture the interplay between the patient and ECMO by continuously collecting perfusion data [15]. Analysis of information-dense perfusion data may be leveraged to improve the prediction accuracy of clinically meaningful outcomes in ECLS care.

Incorporating more granular data requires a new model that is capable of integrating categorical and time-series data. The prevalence of recurrent neural networks (RNNs) in health care data science has increased recently. The ability of RNNs to...
efficiently understand time dependencies makes this approach beneficial in certain types of medical data, such as ventilator settings [16], vital signs [17], medication administration [18], imaging studies [19], and radiology reports [20]. One type of RNN, long short-term memory (LSTM), is specifically designed for long time series, such as our data set with weeks-long hospital courses. LSTM can encode these time series into a compressed latent space, which can be concatenated with static variables, such as age, gender, and other clinical characteristics.

**Novelty**

The innovation of our study is two-fold: (1) data source and (2) algorithm design. Perfusion data were collected from the ECMO devices and recorded at highly granular intervals. Our analysis sheds new light on the effectiveness of ECMO. Second, unlike prior work using laboratory values, clinical scores, and other static data, the patient information used in our study was both dynamic and static. Using a 2-headed neural network, our predictive algorithm efficiently incorporates static information, such as sex and clinical scores, along with dynamic data. LSTM networks encode the perfusion time-series data into a latent space, which is then concatenated with an encoding of the static variables. This new latent space was used to classify patients.

We present the Continuous Evaluation of VV-ECMO Outcomes (CEVVO) predictive model for determining successful decannulation from VV-ECMO using both pre- and postcannulation data. When using both, the model can continuously update its prediction, providing a running measure for patient potential recovery. Such a measure may help clinicians and patient families make more informed decisions about care. Using synthetic data sets, we demonstrate that understanding time dependence is the essential ingredient to accurate predictions. Our framework also guides the categorization of patients into high-risk and low-risk groups, alerting care providers about which patients may be better candidates for weaning trials and decannulation.

**Methods**

**Problem Formulation**

Health care data of this type can be presented in two components: (1) clinical information that remains unchanged over the ECMO course, such as age and sex, which are considered static features, and (2) variables that change over time, such as laboratory values and perfusion data, which are considered temporal variables. This study follows the conventions presented in the study by Yoon et al [21]. We define \( S \) as a vector space of static features, and \( X \) as a vector space of temporal features. Let \( S \in S \) and \( X \in \mathbb{R}^k \) be random vectors with specific values denoted by \( s \) and \( x \). Each patient is a tuple of \((s, x_{1:T})\), where \( T \) is the number of time steps. For clarity, patients in our training set were indexed by \( n \in 1,...,N \). Therefore, the training data set is denoted as \( D = \{s_n, x_{n,1:T}\}^{N}_{n=1} \). Each patient also had a categorical outcome \( y \in \{0,1\} \), which forms vector \( Y \) across all patients, with 0 representing unsuccessful decannulation and 1 representing success. We define the probability distribution \( p(Y|S, X_{1:T}) \), and our goal is to use training data \( D \) to learn a density \( \hat{p}^*(Y|S, X_{1:T}) \) that best approximates \( p(Y|S, X_{1:T}) \). This is achieved through the optimization in equation 1:

\[
\text{Min}_{\pi} \mathcal{D}_{KL}(p(Y|S, X_{1:T}) \parallel \hat{p}^*(Y|S, X_{1:T})) \tag{1}
\]

The abovementioned Kullback-Leibler divergence can be calculated through the loss function in equation 2. This is identical to the cross-entropy because the entropy of the ground truth distribution is 0. The model can best approximate the true distribution by using backpropagation to minimize equation 2:

\[
L = (-1/N) \sum_{n=1}^{N} (y_n \log(\hat{y}_n) + (1-y_n) \log(1-\hat{y}_n)) \tag{2}
\]

**Synthetic Data Set**

We hypothesized that the high performance of the LSTM-based architecture is owing to its superior ability to capture long-term dependencies in the data set. To test this notion, 2 synthetic data sets of size \( N=234 \) and \( T=2054 \) were generated using a Gaussian process regression (GPR) model [22]. As GPR is nonparametric, it can generate synthetic data without making assumptions about the underlying relationships between variables and dynamics over time. By tuning parameters of the generative model, we can adjust the strength of long-term dependencies in the data. Using the GPR model, we sample data from a multivariable normal distribution, in which the covariance encodes dependencies between time points as shown in equation 3:

\[
f(\mu, \Sigma) \tag{3}
\]

where \( f \) denotes the expected values of the inputs and denotes the covariance. The covariance is encoded by a radial basis function (RBF) kernel, as shown in equation 4. The length scale parameter \( L \) of the RBF adjusts the local smoothing. Higher values for this parameter encode dependencies over a longer period, leading to smoother dynamics.

\[
k(x, x') = \exp(-d(x, x')^2/(2L^2)) \tag{4}
\]

where \( d(x, x') \) denotes the Euclidean distance. The long-term dependencies are captured by the probability of observing specific values conditioned on earlier time points. This assumption is reasonable in our application to VV-ECMO and not necessarily held in previous models such as logistic regression and some deep neural networks. The performance of previous models on GPR data is thus not affected by different choices of length scale, whereas the LSTM-based model should lose its advantage with increasing length scale.

Two groups of synthetic data were created: the first with \( L=1 \) and the second with \( L=100 \), and it was expected that CEVVO would be the only one to perform substantially better on \( L=1 \). The other models should have similar performance between \( L=1 \) and \( L=100 \). The length scale had to be larger than or equal to each time step; therefore, \( L=1 \) was close to the minimum allowable length scale.

**Model Design**

A general overview of this framework is presented in Figure 1. It is composed of 2 independent heads: a static (clinical) data encoder and a temporal feature (perfusion) encoder. Theoretically, each distills the relevant information from the 2 data sets (clinical and perfusion) beforeconcatenating them in the classification block.
The static information encoder is based on an autoencoding scheme along with an additional final dense layer. The first dense layer had 32 nodes, the second layer had 33, and the final layer had 25 nodes. These dimensions were chosen via Bayesian optimization hyperparameter tuning implemented through the Keras Python package by Chollet et al [23].

The perfusion information encoder was based on LSTM layers. These recurrent networks were found to work exceedingly well, as they were built on the assumption that earlier time points have marginal effects on later time points. A 1x1 convolutional layer was first used to expand the feature map before the LSTM to create a projection shortcut and act as a filter. The tanh activation function allowed the convolution layer to increase, decrease, or negate certain input values. Although an additional LSTM layer could do this processing, the convolution layer contained significantly fewer parameters. The filter size for the convolution and the LSTMs was 1024, which was also chosen via Bayesian optimization hyperparameter tuning.

The classification block concatenated the final outputs of the clinical information encoder and the perfusion encoder. By this point, the original clinical inputs were reduced from 32 to 25, and the original perfusion inputs were reduced from 16,432 to 1024. These 2 final layers were concatenated into a final layer of 1049. This led to a single output neuron with a sigmoid activation, which acted as the final prediction. This prediction was then compared with the ground truth, and the loss was calculated using the binary cross-entropy. The average of all the losses was calculated with equation 2. These losses were backpropagated through the network to make the probability distribution generated by the model resemble the reality.
Defining Patient Risk Grouping

For the risk groups to have meaning, the calibration of the model must be assessed. A calibration plot for the training set was created and showed an S-shaped misalignment. The misalignment was corrected using Platt scaling.

Four clinical groups were defined with respect to the calibrated mean and SD of the model’s predictions on $D$. Let $M_D$ and $S_D$ be the mean and SD of the sigmoid output values of training data $D$. The grouping was determined according to equation 5:

\[
\begin{align*}
 f_{\text{group}}(x) &= \begin{cases} 
 0, & \text{if } x \leq M_D - S_D \\
 1, & \text{if } M_D - S_D < x \leq M_D \\
 2, & \text{if } M_D < x \leq M_D + S_D \\
 3, & \text{if } M_D + S_D < x
\end{cases}
\end{align*}
\]

Ethical Considerations

This study was conducted in accordance with the institutional review board of the Columbia University (#AAAT0563).

Data

A retrospective chart review was performed, and continuous perfusion data and clinical information were collected from 118 patients cannulated to VV-ECMO at a high-volume ECMO center’s intensive care unit between January 1, 2020, and December 31, 2021. Patients reconfigured to venoarterial-venous or venoarterial were excluded.

Patient data were collected from Spectrum Medical software (Quantum Informatics), which records data from each patient’s ECMO machine. Six relevant perfusion variables were selected with expert insight and were collected at 120-second intervals. These were the pressure change across the membrane lung, the venous drainage pressure, the blood flow across the ECMO circuit, the pump head rotation speed (needed to generate the blood flow), the sweep gas flow (rate of oxygenated gas flowing through the membrane lung), and length of time the patient was supported on ECMO. Two additional perfusion variables were created to account for differences between patients: the flow across the pump divided by the patient’s BMI and the sweep gas flow divided by the flow across the pump. In addition to these 8 perfusion variables, 12 clinical variables were selected: decannulation result, age, sex, cause of respiratory distress, BMI, cardiac arrest before ECMO, shock (ie, hemodynamic instability) before ECMO, reinfusion and drainage cannulation location, reinfusion and drainage cannula size, and the type of ventilation provided (Table S2 in Multimedia Appendix 1). The 12 clinical variables included the outcome label, which was not included in the input data. Further clinical information that was not included in the model can be found in Table S3 in Multimedia Appendix 1. An example of 5 successful and 5 unsuccessful patients is shown in Figure 2. The chaotic nature of the perfusion variables helps to justify more advanced machine learning methods.
To enable incorporation of all time points in each VV-ECMO run, the first preprocessing step involved truncation, which refers to clipping the perfusion data set at different percentages of the total run. For each patient, in addition to the 100% of the ECMO run (ie, the full run), the first 90%, the first 80%..., the first 10% of the run were appended as additional runs. Thus, the full data set involved 1180 sequences of data points, 10 for each patient. Each data point consisted of a 3D perfusion time series (patient deidentified ID code, time step, and variable) and 2D clinical data (patient deidentified ID code and variable). Owing to varying ECMO run lengths, each time-series sequence was standardized to 2054 time steps. This length was the largest size possible, given the GPU constraints. Standardization was performed by averaging dense time steps and forward-filling empty steps. The remaining empty time steps were set to 0. Truncations were treated as full runs, that is, the final values for the 10% and the 100% truncation occurred at the same time step—2053. Each truncation is, in effect, stretched over the 2054 time steps. This ensures that the model is not given hints about which truncation it is seeing.
The performance of the model was evaluated through cross-validation. In each iteration, the list of patient IDs was randomized and split into 5 groups of 23 patients, each with 18 successful and 5 unsuccessful patients. Three random patients were excluded to have 5 groups of the same size. Three groups of patients (69/115, 60% of the total) were chosen as the training data set, one group (23/115, 20%) was chosen as the validation set, and one group (23/115, 20%) was chosen as the test set. This process was repeated 5 times until each group had been included in the test set once. The patient list was then randomized again to begin the next cross-fold validation. This ensured that the training set, validation set, test set, and unused patients differed each time. In total, there were 12 iterations of this 5-fold cross-validation.

Each set of training data consisted of 69 patients, and the validation and test sets had 23 patients. Including all truncations, the training set had 690 data points, and the validation and test sets had 230 data points each.

The data sets were then scaled using MinMaxScaler from the sklearn Python package by Pedregosa et al [24]. The scaler was trained on the training data and then used to transform all 3 sets.

**Synthetic Data Set**

The GPRs were generated using the Gaussian process regressor Python package sklearn Pedregosa et al [24]. Two different data sets were generated with different values for the length scale of the RBF kernel (1, 100). A GPR model was first fit to unnormalized perfusion data from all patients. To generate more realistic synthetic data, patients were divided into successful and unsuccessful decannulation groups and sorted according to the ECMO run time. They were then grouped into triplets based on these criteria, which resulted in 30 successful triplets and 8 unsuccessful triplets. To create the training data for the GPRs, each patient’s age, gender, and BMI were extracted and normalized using the StandardScaler from the sklearn Python package by Pedregosa et al [24]. These, in addition to a time-step value, were treated as independent variables. The dependent variable was the unnormalized perfusion data from the triplet. Each GPR was trained only on a single perfusion variable, so each triplet had 8 GPRs, 1 for each perfusion variable. For each of the 30 successful triplets, each GPR model was sampled 3 times for a total of 90 synthetic successful patients. For the 8 unsuccessful triplets, each GPR was sampled 18 times for a total of 144 synthetic unsuccessful patients. A diagram of this process is shown in Figure 3.

**Model Assessment**

After each model was trained, predictions were calculated for the test sets. Each prediction varied between 0 and 1 owing to

---

**Figure 3.** Diagram of the process of generating synthetic patient data. Solid boxes indicate patient data (including synthetic), and dotted boxes indicate a Gaussian process regression (GPR) model fit to real patient data. The sampling of the GPRs step was repeated 2 times, 1 for each kernel length scale.
the sigmoid activation in the final neuron. To assess performance, area under the receiver operating characteristic curve (AUROC) and the average precision (AP) were calculated using the sklearn package. AP approximated the area under the precision-recall curve. The predictions and ground truths were sampled 5000 times with replacement to create the AUROC and AP CIs. The bootstrapping pseudocode for estimating the AUROC CIs can be found in algorithm S1 in Multimedia Appendix 1. This bootstrapping code was then repeated for different subsets of the data. The AUROC and AP CIs were calculated for each day after cannulation between 0 and 24 (e.g., the AUROC and AP for all data points ending on day 10, including truncations). These values were plotted along with their CIs. For the synthetic data, the bootstrapping method was only used on the entirety of each data set.

A successful model is expected to provide accurate and reliable insight into whether a patient will be decannulated.

**Results**

**Model Performance on Real Data**

The model’s overall performance on the real data achieved an average AUROC of 0.6937 (95% CI 0.6822-0.7055). The mean AP was 0.8599 (95% CI 0.8515-0.8682).

A clinically relevant breakdown is AUROC and AP by day, as shown in Figures 4 and 5. Therefore, we observed that tight CIs begin to expand after day 11 as the number of data points decreased. By limiting the time frame to only include patient data points sampled between days 3 and 11, the AUROC 95% CI was 0.7048-0.7428, and the AP 95% CI was 0.9074-0.9261.

*Figure 4.* The area under the receiver operating characteristic curve (AUROC; in green) computed from all samples within a 1-day time frame, for example. AUROC for samples collected between days 0 and 1 are shown on day 1. Purple bars indicate the number of data points occurring on that day (right y-axis).

*Figure 5.* The average precision (AP; in green) computed from all samples within a 1-day time frame, for example. AP for samples collected between days 0 and 1 are shown on day 1. Purple bars indicate the number of data points occurring on that day (right y-axis).

**Model Comparison on Real Data**

As detailed in Table S2 in Multimedia Appendix 1, ECMOnet, Predicting Death for Severe Ards on VV-ECMO, Respiratory ECMO Survival Prediction, Roch, Venovenous ecmo mortality score, and Prediction of Survival on ECMO Therapy score rely on either logistic regression or recursive partitioning analysis to determine the patient grouping or scoring classification. To provide a fair comparison with the proposed model, the AUROC and AP calculations were repeated with a logistic regression.
model and a decision tree. Both models were trained on the same training data and assessed on the same test data as the proposed model. Moreover, the 95% CIs were determined with the bootstrapping algorithm presented in algorithm S1 in Multimedia Appendix 1. Furthermore, to provide a comparison with the study by Ayers et al [13], a dense neural network was included. Finally, a Naive Bayes model was included to demonstrate the necessity of including dependence between time points. Unlike the LSTM, Naive Bayes assumes conditional independence between features, making previous models incapable of understanding the time series as anything beyond a bag of values. Table 1 demonstrates that CEVVO is the most effective model for ECMO data, showing a significant improvement compared with other methods. Using a permutation test, CEVVO demonstrated a significantly higher AUROC (all \( P \) values <.001) and AP (all \( P \) values <.04) than all other methods.

Table 1. Comparison of Continuous Evaluation of Venovenous Extracorporeal Membrane Oxygenation Outcomes (CEVVO) with other models used previously.

<table>
<thead>
<tr>
<th>Model name</th>
<th>Total AUROC(^a), 95% CI</th>
<th>Total AP(^b), 95% CI</th>
<th>( P ) value compared with CEVVO (AUROC)</th>
<th>( P ) value compared with CEVVO (AP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEVVO</td>
<td>0.6822-0.7055</td>
<td>0.8515-0.8682</td>
<td>&lt;.001</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Logistic regression</td>
<td>0.6395-0.6626</td>
<td>0.8396-0.8566</td>
<td>&lt;.001</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Naive Bayes</td>
<td>0.5876-0.6081</td>
<td>0.8111-0.8255</td>
<td>&lt;.001</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Dense network</td>
<td>0.5673-0.5908</td>
<td>0.8148-0.8322</td>
<td>&lt;.001</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Decision tree</td>
<td>0.5419-0.5596</td>
<td>0.5273-0.5467</td>
<td>&lt;.001</td>
<td>&lt;.001</td>
</tr>
</tbody>
</table>

\(^a\)AUROC: area under the receiver operator characteristic.

\(^b\)AP: average precision.

\(^c\)Not applicable.

**Risk Classification System**

The calibration plot of the training data is shown in Figure S1 in Multimedia Appendix 1. The classic S-shaped misalignment indicated that Platt scaling would improve the calibration. Both the calibrated training and test sets are shown in Figure 6.

Using the predictions as an indication of favorable or unfavorable outcomes, patients can be stratified into groups based on their prediction value using equation 5. The clinically relevant measures of performance are shown in Figures 7 and 8. These charts were created by finding the nearest predicted value of each patient before either 72 or 96 hours, sorting them into groups according to equation 5, and then charting their decannulation result. Patients decannulated before 72 or 96 hours were excluded. In the 72-hour case, the groups had a successful decannulation rate of 58% (7/12) for group 0, 77% (17/22) for group 1, 88% (42/48) for group 2, and 92% (11/12) for group 3. In the 96-hour case, the groups had successful decannulation rates of 54% (6/11), 85% (17/20), 81% (42/50), and 100% (9/9), respectively.

**Figure 6.** The calibration plot for both the training and test set prediction. Each set of predictions has been scaled. The green line shows the theoretical perfect calibration, and the purple bars show the number of data points in each bin.
A Boschloo exact test between groups 0 and 3 yielded $P$ values of .04 for 72 hours and .01 for 96 hours.

**Necessity of Time Dependencies**

To ensure that each of the synthetic data sets were comparable with each other and the original, a $t$-distributed Stochastic Neighbor Embedding [25] was used (Figure 9). A more concrete example is shown in Figure 10, where a single synthetic input was run through both the $L=1$ and $L=100$ GPRs and then compared with an original patient.

The procedure specified in the Model Assessment section was repeated for CEVVO, logistic regression, and dense network on the synthetic data set. The 95% CI for the AUROC is shown in Table 2. The expected result is observed where logistic regression and the dense network show no change in performance. CEVVO shows a significant drop in performance.
despite having similar, nonlinear properties to the dense network.

**Figure 9.** 2D t-distributed Stochastic Neighbor Embedding (tSNE) of the 2 synthetic sequential data set and the original patient data. Each dot represents a synthetic patient; the red dots indicate data generated using a radial basis function (RBF) with $L=1$, the green dots indicate data generated using length $L=100$, and the blue dots indicate the original data. The significant overlap connotes similarity between the literal values.

**Figure 10.** An example synthetic patient, shown in both the $L=1$ and $L=100$ data sets compared with a similar real patient (in blue). ECMO: extracorporeal membrane oxygenation; RPM: revolutions per minute.

**Table 2.** Comparison of Continuous Evaluation of Venovenous Extracorporeal Membrane Oxygenation Outcomes (CEVVO) with top-performing models used previously on each synthetic data set.

<table>
<thead>
<tr>
<th>Model name</th>
<th>Total AUROC\textsuperscript{a} for $L=1$ synthetic data set, 95% CI</th>
<th>Total AUROC for $L=100$ synthetic data set, 95% CI</th>
<th>$P$ value between $L=1$ and $L=100$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEVVO</td>
<td>0.8223-0.8583</td>
<td>0.7424-0.7849</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Logistic regression</td>
<td>0.7813-0.8213</td>
<td>0.7814-0.8190</td>
<td>.46</td>
</tr>
<tr>
<td>Dense network</td>
<td>0.7080-0.7513</td>
<td>0.6924-0.7352</td>
<td>.17</td>
</tr>
</tbody>
</table>

\textsuperscript{a}AUROC: area under the receiver operator characteristic.
Discussion

Principal Findings

VV-ECMO is an invasive and resource-intensive therapy used for patients with refractive respiratory failure. Decannulation from ECMO is generally performed through a weaning trial, in which the ECMO support, measured as flow through the circuit, is titrated down. Experienced clinician decision-making with careful consideration of patient hemodynamics, response to therapy, and pathophysiology informs the decision on when to perform the weaning trial. Our study investigates a novel approach to analyzing clinical information and perfusion hemodynamics in real time to assist clinicians with the decision of when to move forward with decannulation from VV-ECMO.

Although CEVVO was more accurate at predicting the success of decannulation than other models, the model should be considered as an additional data point to guide clinical management. Patients stratified to the high-risk group had a higher risk of therapy failure, with >50% of the patients in this group successfully decannulated in both the 72 and 96 hour cases. As expected, the calibration plot also showed that patients in the low-risk group were decannulated successfully more often. Using these data, clinicians may reference the model and elect to start weaning trials on patients stratified to the low-risk cohort sooner.

Comparison With Prior Work

To the best of our knowledge, CEVVO is the first to use ECMO perfusion data and a deep learning architecture to provide clinical decision support for the decannulation decision for VV-ECMO. By using a model that can successfully combine dynamic and static data, significantly improved performance on binary classification can be achieved when compared with other models. Using perfusion data and clinical information, CEVVO was trained to classify patients by decannulation outcome (successful or unsuccessful). The performance was evaluated using 3 criteria: AUROC, AP, and the clinical usefulness of predictions. Relative to other models noted in the literature, such as logistic regression and decision trees, the LSTM-based model showed significant improvement on the ECMO machine data set.

Performance

The AUROC and AP scores for the full data set had 95% CIs of 0.6822-0.7055 and 0.8515-0.8682, demonstrating a fair ability to predict exact outcomes. This was marginally improved to 95% CIs of 0.7048-0.7428 and 0.9074-0.9261 by limiting the data set to only consider data points collected 3 to 11 days after cannulation. However, these numbers only represented the average performance.

Synthetic Data

The use of GPR-created data sets further cemented the notion that the novelty of the architecture, understanding time dependence, is truly what is responsible for the performance edge over other models. The assumption of temporal dependence is inherent in the data as it is medically motivated. There is an expectation that the specific value of the perfusion data shares much mutual information with the outcome. The L=1, L=100, and original data sets are very similar in their t-distributed Stochastic Neighbor Embedding projection, differing only slightly in the specific values. However, as shown in Figure 10, within the L=100 group, the local structure was obliterated, leading to a loss of information about how later time points affect the outcome probability. Logistic regression explicitly assumes that each time point is independent, and thus, it has highly similar AUROC distributions ($P<.46$) compared with the dense network ($P=.17$) and CEVVO ($P<.001$). The nonlinear nature of dense neural networks is able to approximate time dependence but is less efficient than the LSTM-based architecture.

Risk Classification

These initial measures of performance were then used to contextualize the clinical predictions: stratifying people into groups, based on associated risk, to predict recovery. The numerical value of each patient’s prediction was divided into groups, and patients were followed to their decannulation result. For the grouping to be useful, there should be some difference in the success percentage that increases from the high-risk group to the low-risk group. This result was observed in this study. When measured at 72 hours, 58% (7/12) of the patients in the high-risk group had a successful decannulation, whereas 92% (11/12) of the patients in the low-risk group were successfully decannulated ($P=.04$). When measured at 96 hours, the successful decannulation percentage was similar: 54% (6/11) of the patients in the high-risk group and 100% (9/9) patients in the low-risk group were successfully decannulated ($P=.01$).

Limitations

Cohort studies using retrospective data collection are subject to inherent bias. We mitigated bias in this study by including all consecutive patients supported on VV-ECMO at our center. Incomplete data recording from the ECMO devices may have contributed to this model. In the future, this could be mitigated by increasing the sample size and improving data capture methodology.

 Clinically, patients with different indications for ECMO support vary in their hospital course, and the number of different disease etiologies may have been too few for the model to learn. Larger cohorts may help mitigate the issues related to an underpowered data set. Furthermore, model performance declined beyond a period of approximately 11 days, which may be attributed to a challenging hospital course with heterogeneous factors and an increased risk for complications. The effectiveness of ECMO as a long-term therapy remains unclear, and our data support this conclusion.

Future Direction

In the future, more information about each patient’s hospital course, such as administration of vasopressors, ventilator settings, imaging studies, and other interventions may be used to develop an improved model. Indeed, more data and reducing unaccounted variables may improve model performance over longer periods. Extending this study to include patients on other forms of ECLS, such as VA-ECMO and cardiogenic shock, respectively.
may be helpful in guiding clinical management. We suggest that larger and more comprehensive repositories of health care data may improve the management of patients considered most critically ill.

Acknowledgments
The authors would like to thank members of the Columbia University Irving Institute for Cancer Dynamics, Data Science Institute, the Biomedical Engineering Department, and the Columbia University Irving Medical Center for helpful discussions and support.

Data Availability
The main clinical data set is unavailable for the public because it contains protected health information. However, 2 synthetic data sets generated based on the original data are available at the Aizilab GitHub data set, along with the corresponding Python code.

Authors' Contributions
JF played a key role in conceptualization and data curation and took the lead in formal analysis, investigation, methodology, software, visualization, and writing the original draft. AA led data curation and provided support in conceptualization, investigation, and writing the review. DM and JB contributed to data curation. PL led conceptualization, supported investigation, and co-led supervision, contributing to the writing of the review. EA provided support in conceptualization, formal analysis, investigation, methodology, and software. In addition, EA co-led supervision and contributed to writing the review.

Conflicts of Interest
None declared.

Multimedia Appendix 1
Demographic data for the patient population, along with the bootstrapping algorithm.

[DOCX File, 79 KB - biomedeng_v9i1e48497_app1.docx ]

References


Abbreviations

AUROC: area under the receiver operating characteristic curve
AP: average precision
CEVO: Continuous Evaluation of Venoarterial Extracorporeal Membrane Oxygenation Outcomes
ECLS: extracorporeal life support
ECMO: extracorporeal membrane oxygenation
GPR: Gaussian process regression
LSTM: long short-term memory
RBF: radial basis function
RNN: recurrent neural network
VA-ECMO: venoarterial extracorporeal membrane oxygenation
VV-ECMO: venovenous extracorporeal membrane oxygenation
Joshua Fuller, Alexey Abramov, Dana Mullin, James Beck, Philippe Lemaitre, Elham Azizi. Originally published in JMIR Biomedical Engineering (http://biomedeng.jmir.org), 02.02.2024. This is an open-access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work, first published in JMIR Biomedical Engineering, is properly cited. The complete bibliographic information, a link to the original publication on https://biomedeng.jmir.org/, as well as this copyright and license information must be included.
Enhancing Energy Efficiency in Telehealth Internet of Things Systems Through Fog and Cloud Computing Integration: Simulation Study

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Abstract

Background: The increasing adoption of telehealth Internet of Things (IoT) devices in health care informatics has led to concerns about energy use and data processing efficiency.

Objective: This paper introduces an innovative model that integrates telehealth IoT devices with a fog and cloud computing–based platform, aiming to enhance energy efficiency in telehealth IoT systems.

Methods: The proposed model incorporates adaptive energy-saving strategies, localized fog nodes, and a hybrid cloud infrastructure. Simulation analyses were conducted to assess the model’s effectiveness in reducing energy consumption and enhancing data processing efficiency.

Results: Simulation results demonstrated significant energy savings, with a 2% reduction in energy consumption achieved through adaptive energy-saving strategies. The sample size for the simulation was 10-40, providing statistical robustness to the findings.

Conclusions: The proposed model successfully addresses energy and data processing challenges in telehealth IoT scenarios. By integrating fog computing for local processing and a hybrid cloud infrastructure, substantial energy savings are achieved. Ongoing research will focus on refining the energy conservation model and exploring additional functional enhancements for broader applicability in health care and industrial contexts.

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KEYWORDS
cloud computing; energy-efficient; fog computing; Internet of Things; IoT; telehealth

Introduction

Overview
Health care is a critical global industry, and the advent of the Internet of Things (IoT) and cloud computing has significantly transformed health care system management. The ever-increasing data volume generated by these systems demands efficient, energy-saving computing platforms. In response, we present a groundbreaking energy-efficient model that seamlessly integrates telehealth IoT devices with fog and cloud computing–based platforms, offering a unique solution to address energy efficiency and data processing challenges. The rapid proliferation of IoT devices in health care has transformed approaches to patient care, diagnostics, and treatment. Telehealth, a key IoT health care application, has proven its potential to enhance care quality, reduce costs, and boost patient satisfaction. Despite these benefits, issues such as scalability, latency, and resource management persist, along with the significant challenge of energy consumption in smart devices within fog environments [1]. As a result, energy efficiency must be prioritized in the development of fog computing solutions, given its substantial impact on reducing carbon footprints and mitigating climate change effects. The
large-scale deployment of telehealth IoT devices also raises concerns about energy consumption and data processing efficiency in delivering quality health care services. Intelligent choices for telehealth IoT devices should consider factors such as device movement or relevant environmental conditions to optimize energy consumption and manage associated equipment effectively. Typically, cloud-based analytical assessments are conducted for these devices [2]. To tackle these challenges, we propose an energy-saving model that integrates telehealth IoT devices with a fog and public or private cloud computing–based platform. The aim of the study is to develop an energy-efficient model that optimally integrates telehealth IoT devices with fog and cloud computing platforms, addressing challenges related to energy consumption, scalability, and data processing efficiency in delivering quality medical and patient services.

Telehealth IoT devices refer to a wide range of interconnected medical devices and sensors that facilitate remote health care services. These devices enable the continuous monitoring of patient’s vital signs, timely diagnostics, and personalized treatment plans, thereby improving the overall quality of health care. Some common examples of telehealth IoT devices include wearable health monitors, smart glucose meters, remote patient monitoring systems, and telemedicine platforms. The large-scale deployment of telehealth IoT devices presents several challenges [3], including energy consumption, data management, latency, security and privacy, scalability, and interoperability.

Related Work

Telehealth has emerged as a promising solution to address various challenges in health care, such as accessibility, cost, and quality of care [4]. IoT devices play a significant role in telehealth applications, enabling remote monitoring, diagnostics, and treatment [2]. Several studies have investigated the implementation and efficacy of telehealth IoT devices in various health care scenarios, highlighting their potential to improve patient outcomes and satisfaction [5,6]. Fog computing has been identified as a promising approach to address the challenges associated with large-scale IoT deployments in health care, such as latency, energy consumption, and data management [7,8]. Researchers have proposed several fog computing–based architectures and frameworks for health care applications, demonstrating the potential of fog computing to enhance the performance and efficiency of telehealth IoT devices [9-11].

Cloud computing has gained significant attention in health care due to its scalability, cost-effectiveness, and advanced data analytics capabilities [12,13]. Several studies have explored the integration of cloud computing with telehealth IoT devices, showing its potential to address the challenges related to data storage, processing, and security [14-16].

Energy efficiency is critical in large-scale IoT deployments, especially in health care applications where device longevity and reliability are essential [17]. Researchers have proposed various energy-saving models and strategies for IoT devices, including adaptive power management [18], energy-efficient routing protocols [19], and data compression techniques [20]. However, few studies have specifically focused on energy-saving models that integrate telehealth IoT devices with fog and cloud computing–based platforms. The integration of fog and cloud computing has emerged as a promising approach to harness the benefits of both paradigms and address the challenges of large-scale IoT deployments [21,22]. Several studies have proposed models and frameworks that combine fog and cloud computing for various IoT applications [23-25], but few have specifically targeted energy-saving in telehealth IoT deployments.

In recent years, several simulation methods have been developed to study the integration of fog nodes in IoT devices and cloud computing. Gupta et al [26] introduced iFogSim, a toolkit for modeling and simulating resource management techniques in IoT, edge, and fog computing environments. Oueis et al [27] presented a simulation study on load distribution in small-cell cloud computing using fog computing and proposed a fog balancing technique to optimize resource allocation and reduce latency. Barcelo et al [28] explored IoT-cloud service optimization through simulation in smart environments, presenting a novel optimization framework that uses fog nodes to reduce latency and energy consumption. Zeng et al [29] conducted a comparative study of IoT cloud and fog computing simulations using iFogSim and Cooja, discussing the advantages and limitations of both simulators and providing insights into selecting an appropriate tool for specific scenarios. Lastly, Byers and Wetterwald [30] discussed the concept of fog computing and its importance in distributing data and intelligence for IoT resiliency and scalability, presenting various simulation models and techniques used to evaluate the performance of fog computing in IoT environments. Several studies have focused on the Yet Another Fog Simulator (YAFS) framework, a simulator designed for modeling and simulating fog computing environments in IoT scenarios. Bermejo et al [31] introduced YAFS, presenting the architecture, components, and use cases of the simulator, demonstrating its effectiveness in modeling and simulating fog computing deployments. García et al [32] showcased YAFS’s ability to model and simulate fog computing scenarios and analyze the performance of different scheduling algorithms. In a comparative study, Rodríguez et al [33] analyzed the features, capabilities, and limitations of YAFS, iFogSim, and EdgeCloudSim simulators, providing insights into selecting the most suitable tool for specific fog computing scenarios.

Several studies have explored different aspects of telehealth simulations, fog nodes, IoT devices, and cloud computing for energy-saving purposes. Aazam and Huh [34] discussed a smart gateway–based communication approach using fog computing for energy-saving in the Cloud of Things, which can be applied to various IoT applications, including telehealth. Verma and Sood [35] presented a fog-assisted IoT framework for patient health monitoring in smart homes, focusing on energy efficiency and reduced latency through a decentralized fog computing architecture. Koubaa et al [36] proposed a fog-based emergency and health care system for smart cities, which leverages fog nodes and IoT devices to optimize energy consumption and provide real-time health care services, thus addressing energy-saving concerns in telehealth scenarios. Sareen et al [37] introduced an energy-efficient context-aware framework for managing application execution in cloud-fog environments.
which can potentially improve energy efficiency in various IoT applications, including telehealth scenarios.

**Methods**

**Model Overview**

The proposed energy-saving model is designed to integrate telehealth IoT devices with a fog and cloud computing–based platform, leveraging the advantages of both paradigms to optimize energy consumption and ensure efficient data processing. The model comprises 3 main components: IoT devices, fog nodes, and public or private cloud servers, which are interconnected through a communication network.

The model architecture is shown in Figure 1 [38].

1. **IoT devices**: telehealth IoT devices, such as wearables, sensors, and remote monitoring systems, collect and transmit patient data in real time. These devices can dynamically adjust their power states (eg, active, idle, and sleep) based on their tasks, reducing energy consumption without compromising the quality of health care services.

2. **Fog nodes**: fog nodes, located near IoT devices, serve as intermediate processing units. They perform localized data processing, analytics, and storage, reducing the amount of data transmitted to the cloud servers.

3. **Cloud servers**: cloud servers provide a robust infrastructure for large-scale data storage, processing, and advanced analytics.

4. **Communication network**: a communication network connects IoT devices, fog nodes, and cloud servers, enabling seamless data transmission and task allocation.

Figure 1. Telehealth Internet of Things (IoT) devices integrated with fog nodes and a private or public cloud architecture model. LAN: local area network.

The telehealth IoT network depicted in the diagram is designed to ensure efficient and secure data transmission between the different network components. To ensure network security, firewalls are placed between IoT devices and fog nodes. This ensures that unauthorized access to the network is prevented, and sensitive health care data are kept confidential. To process the data requests, the fog nodes are equipped with data analytics functions that enable them to intelligently assign different types of requests to either fog nodes, a private cloud, or a public cloud. This intelligent decision-making process is more effective and efficient than the traditional “first-come, first-served” approach. The gateway and router are integral components in the network that enable seamless data transmission between the fog nodes and cloud instances. The gateway acts as the entry point for the network and connects the IoT devices to the local fog nodes. It is responsible for handling the data transmission and conversion between different protocols used by IoT devices and fog nodes. The router, on the other hand, is responsible for directing the data traffic between the fog nodes and cloud instances based on various factors, such as the sensitivity, priority, and latency requirements of the data. It determines which data should be sent to the cloud and which data should be processed by the fog nodes, ensuring efficient use of network resources. The router also handles the communication between different fog nodes and cloud instances, enabling seamless data transmission across the network.

The proposed telehealth IoT system shown in Figure 2 intelligently manages data transmission based on the sensitivity and priority of the data. For high-sensitivity data, the system ensures privacy and security by sending it directly to the private cloud, which then transfers the data to authorized health facilities as needed. On the other hand, low-sensitivity but high-priority requests are routed to the fog nodes as they have the capability to process urgent requests in a timely manner, such as in life-threatening emergency situations. These requests are then transmitted to ambulance systems for immediate treatment. Lastly, data with low sensitivity and low priority are sent to the...
public cloud as it has more space and scalability to store and process such data. The public cloud can also serve as a repository for future research or clinical purposes.

By allocating data transmission to the appropriate destination, the proposed system ensures efficient and effective data processing while maintaining privacy and security for sensitive health care data. This approach also optimizes energy consumption and reduces latency, ensuring a seamless experience for health care providers and patients. The categorization of high and low sensitivity and high and low priority data sent from telehealth IoT monitor devices can depend on various factors, including the specific use case, regulatory requirements, and patient needs. One possible approach could be to use threshold values based on vital signs such as pulse and heartbeat to categorize the data. For example, data related to vital signs that fall within normal ranges may be classified as low sensitivity and low priority, as they do not require immediate attention. Data related to vital signs that are outside the normal range but do not pose an immediate threat to the patient’s health may be classified as low sensitivity but high priority. Data related to vital signs that indicate a life-threatening condition, such as cardiac arrest, may be classified as high sensitivity and high priority, requiring immediate attention from health care providers.

The exact vital sign thresholds for patient emergencies can vary depending on a range of factors, including the age and health condition of the patient, the specific symptoms, and other medical history [39]. In general, some common vital sign thresholds used to classify emergencies include the following:

- Heart rate: a heart rate above 100 bpm or below 60 bpm may be indicative of an emergency [40].
- Blood pressure: a systolic blood pressure (the top number) above 180 mm Hg or below 90 mm Hg, or a diastolic blood pressure (the bottom number) above 110 mm Hg or below 60 mm Hg may indicate an emergency [41].
- Respiratory rate: a respiratory rate above 30 breaths per minute or below 10 breaths per minute may be indicative of an emergency [42].
- Oxygen saturation: an oxygen saturation level below 90% may be indicative of an emergency [43].

However, it is important to note that this is just one possible approach, and the categorization of data should be customized based on the specific needs of the patient and health care provider. It is also important to comply with relevant regulations and ensure patient privacy and security while handling sensitive health care data.

**Figure 2.** Network topology for the proposed Internet of Things (IoT) devices integrated with fog nodes and cloud. A brief overview of the components in the network topology: (1) IoT devices (blue circles) represent individual IoT devices in the network, each associated with a specific fog node. (2) Gateways (GT; orange hexagons) are used to connect the IoT devices to the fog nodes. (3) Fog nodes (FN; green triangles) are intermediate computing resources that process and store data from IoT devices. (4) A router (red square) connects the fog nodes to the private cloud and public cloud. (5) A private cloud (purple square) and a public cloud (yellow square) are the 2 cloud resources in the network.

**Key Components and Energy-Saving Strategies**

The proposed energy-saving model incorporates several strategies to minimize energy consumption.

**Task Allocation**

The model intelligently allocates tasks between fog nodes and cloud servers based on factors such as computational capacity, proximity to IoT devices, and current workload. This ensures
efficient data processing and reduces energy consumption for data transmission.

**Adaptive Power Management**

IoT devices and fog nodes can dynamically adjust their power states (e.g., active, idle, and sleep) based on their tasks and workload, ensuring optimal energy consumption without compromising the quality of health care services.

**Data Compression and Aggregation**

Data generated by IoT devices can be compressed and aggregated at the fog nodes before transmission to cloud servers, reducing the volume of data transmitted and, consequently, energy consumption.

**Network Optimization**

The communication network can be optimized to minimize energy consumption by using energy-efficient routing protocols and minimizing transmission distances.

**Simulation Study**

To assess the effectiveness of the proposed energy-efficient model, we developed a simulation model that emulates a real-world telehealth scenario focused on remote patient monitoring. Within this simulated scenario, numerous patients with chronic conditions are equipped with wearable IoT devices that continuously track vital signs such as heart rate, blood pressure, and blood glucose levels. The gathered data are processed and analyzed by the integrated fog and cloud computing–based platform, facilitating timely diagnostics and personalized treatment plans. **Textbox I** contains the pseudocode for the provided code.

In short, this code is devised to emulate an IoT network, scrutinizing the influence of fog nodes on energy consumption while providing a graphical representation of the network architecture to elucidate the connections among IoT devices, fog nodes, and cloud servers. IoT devices transmit data to their corresponding destinations, such as fog nodes, private clouds, or public clouds, contingent upon their sensitivity and priority attributes. The energy expenditure for data transmission to these target locations differs; hence, the code performs a simulation to determine the residual energy for each device under 2 distinct scenarios (i.e., with and without fog nodes). Subsequently, the code generates a bar chart to depict the energy consumption patterns of IoT devices in both cases, and it stores the energy usage outcomes in 2 separate Microsoft Excel (Microsoft Corporation) files, enabling in-depth examination and assessment of the results.

The algorithm of the code can be analyzed in the following steps:

1. **Initialization:** create IoT devices, fog nodes, and cloud instances with their respective properties.
2. **Connection:** connect IoT devices to fog nodes and then fog nodes, and determine which data are transferred to cloud instances (private and public). Each device is connected to a corresponding fog node.
3. **Data transmission simulation:** simulate data transmission from IoT devices to their respective fog nodes, and then fog nodes assign the requests to a private cloud or a public cloud based on their priority and sensitivity. If the sensitivity of the device is “high,” data are sent to the private cloud. If the sensitivity is “low” and the priority is “high,” there is a chance (defined by `self.fog_node[chance]`) that data are sent to the fog node. If this condition is not met, the device does not send data. If the sensitivity is “low” and the priority is “low,” data are sent to the public cloud.
4. **Energy consumption calculation:** calculate the energy consumed by each IoT device during data transmission, considering the parameter of latency. Different energy costs are associated with sending data to different destinations (fog nodes, private cloud, or public cloud).
5. **Comparison:** compare the energy consumption of IoT devices when using fog nodes and when not using fog nodes. (1) Run the simulation with fog nodes connected and store the remaining energy for each device. (2) Reset the energy of the devices, disconnect them from fog nodes, and run the simulation without fog nodes, storing the remaining energy for each device again.
6. **Export the energy usage results to Excel files for both cases (with and without fog nodes).**
7. **Visualize the network topology with devices, fog nodes, and clouds using the `show_topology` function.**

In this enhanced task allocation algorithm, we incorporate additional factors such as device distance, data sensitivity, request priority, energy consumption, and latency to provide a more sophisticated and adaptable solution for large-scale telehealth IoT deployments. The algorithm starts by defining parameters such as latency, distance, energy consumption, and sensitivity thresholds. The task queues for each fog node and cloud server are initialized. For each task type, average processing times, energy consumption, sensitivity, and priority are calculated for each fog node and cloud server according to some random data sent from each IoT device. The algorithm then assesses the latency, priority, sensitivity, and energy consumption for transmitting data from each device to each fog node and then to the private and public cloud server. Based on these factors, the algorithm selects the optimal fog node and cloud server for each device, ensuring that the chosen nodes meet the specified thresholds for latency, sensitivity, and energy consumption. Tasks are allocated to fog nodes and cloud servers based on data sensitivity, priority, and energy consumption, ensuring that the selected nodes do not exceed the energy consumption threshold. If no suitable nodes are found, alternative energy-saving strategies may be considered, or the energy consumption threshold may be adjusted. Finally, the tasks are processed in fog nodes and cloud servers based on their queues. By considering these additional factors, the enhanced algorithm can provide better energy-saving performance and adaptability to various telehealth scenarios, ensuring that the large-scale deployment of telehealth IoT devices on a fog and cloud computing–based platform remains efficient and effective.

**Textbox 2** contains a task allocation algorithm for telehealth IoT devices integrated with a fog and cloud computing–based platform.
Textbox 1. The pseudocode for the provided code.

1. Define `IoTDevice` class
   - Initialize with attributes: `id`, `distance`, `priority`, `sensitivity`, `fog_node`, `private_cloud`, `public_cloud`, `energy`, `transmit_power`, `idle_power`, and `transmit_time`
   - Define `send_data` method
     - Check if the device has energy left
     - Send high-sensitivity data to private cloud if sensitivity is high
     - Send low-sensitivity, high-priority data to fog node if priority is high and `fog_node` exists
     - Send low-sensitivity, low-priority data to public cloud otherwise
   - Define `idle` method to reduce energy based on idle power and time

2. Define `FogNode` class
   - Initialize with attributes: `id`, `public_cloud`, `energy`, `latency`, `devices`, `fog_energy_cost`, `cloud_energy_cost`, `chance`, `process_power`, `idle_power`, and `process_time`
   - Define `connect_device` method to connect a device to the fog node
   - Define `store_data` method to store data from a device with given sensitivity and priority
   - Define `idle` method to reduce energy based on idle power and time
   - Define `send_data` method to send data from connected devices based on their sensitivity and priority

3. Define `PublicCloud` class
   - Initialize with attributes: `id`, `energy`, `latency`, and `cloud_energy_cost`
   - Define `store_data` method to store data from a device

4. Define `simulate` function
   - Create Internet of Things (IoT) devices with random priority and sensitivity
   - Create fog nodes connected to a public cloud
   - Connect IoT devices to fog nodes
   - Connect IoT devices to private and public clouds
   - Initialize lists to store energy usage results
   - Simulate data transmission with fog nodes, store energy usage results
   - Store energy usage with fog nodes
   - Reset device energy
   - Disconnect devices from fog nodes
   - Simulate data transmission without fog nodes, store energy usage results
   - Store energy usage without fog nodes
   - Create energy usage bar plot and save as an image
   - Save energy usage results to Microsoft Excel files (with and without fog nodes)
1. Define parameters
- Internet of Things (IoT) devices: \( D = \{ d1, d2, ..., dn \} \)
- Fog nodes: \( F = \{ f1, f2, ..., fm \} \)
- Cloud servers: \( C = \{ c1, c2 \} \)
- Task types: \( T = \{ t1, t2, ..., tq \} \)
- Data sensitivity threshold: \( S_t \)
- Data priority threshold: \( Pr_t \)
- Latency threshold: \( L_t \)
- Energy consumption threshold: \( E_t \)

2. Initialize task queues for each IoT device, fog node, and cloud server
- \( Q_D[i] = \{ \} \) for all \( i \) in \( D \)
- \( Q_F[j] = \{ \} \) for all \( j \) in \( F \)
- \( Q_C[l] = \{ \} \) for all \( l \) in \( C \)

3. For each task type \( t \) in \( T \)
   - Calculate the average processing time \( P_t \) and energy consumption \( E_t \) for each IoT device \( i \) in \( D \) and fog node \( j \) in \( F \).
   - Calculate average energy consumption \( E_t \), sensitivity \( S_t \), and priority \( Pr_t \) for each IoT device \( i \) in \( D \) and fog node \( j \) in \( F \).

4. For each device \( d \) in \( D \) and task type \( t \) in \( T \)
   - Calculate the latency \( L_dt \) for transmitting data from device \( d \) to each fog node \( i \) in \( F \) and cloud server \( j \) in \( C \).
   - Calculate the priority \( Pr_{dt} \), sensitivity \( S_{dt} \), energy consumption \( E_{dt} \) for device \( d \) and each fog node \( i \) in \( F \) and cloud server \( j \) in \( C \).
   - Find the fog node \( j^* \) and cloud server \( l^* \) with the minimum latency for device \( d^* \), considering \( Pr_{dt}, S_{dt}, \) and \( E_{dt} \):
     - \( j^* = \text{argmin}_j(L_{dt}) \) for \( j \) in \( F \), such that \( L_{dt} \leq L_t, Pr_{dt} \leq Pr_t \) and \( S_{dt} \leq S_t \)
     - \( l^* = \text{argmin}_l(E_{dt}) \) for \( l \) in \( C \), such that \( L_{dt} \leq L_t, Pr_{dt} \leq Pr_t \) and \( S_{dt} \leq S_t \)

5. Allocate tasks from devices to fog nodes and cloud servers: for each device \( d \) in \( D \) and task type \( t \) in \( T \)
   - If \( S_{dt}[j^*] \leq S_t \), then allocate task \( t \) to cloud server \( l^* \) and add it to the queue: \( Q_C[l^*].append((d, t)) \)
   - If \( Pr_{dt}[j^*] \leq Pr_t \), then allocate task \( t \) to fog node \( j^* \) and add it to the queue: \( Q_F[j^*].append((d, t)) \)
   - Else if \( Pr_{dt}[l^*] \leq Pr_t \), then allocate task \( t \) to cloud server \( l^* \) and add it to the queue: \( Q_C[l^*].append((d, t)) \)
   - Otherwise, consider alternative energy-saving strategies or adjust the energy consumption threshold \( E_t \).

6. Process tasks in fog nodes and cloud servers based on their queues
   - For each fog node \( j \) in \( F \), process tasks in \( Q_F[j] \)
   - For each cloud server \( l \) in \( C \), process tasks in \( Q_C[l] \)

This algorithm aims to balance the load between fog nodes and cloud servers while considering latency, sensitivity, request priority, and energy consumption constraints. It can be further optimized by incorporating additional factors, such as device mobility. It is mainly focused on simulating data transmission from IoT devices to different destinations based on their priority and sensitivity, as well as comparing the energy consumption given the various latency when using fog nodes versus not using them. The objective is to demonstrate the potential benefits of using fog nodes in terms of energy efficiency for IoT devices.

Results

Parameters in Results
Based on the simulation results, we can analyze the impact of different parameters on the energy efficiency and performance of the proposed telehealth model with and without fog computing. The parameters in the results are given below.

Snapshot Interval
The snapshot interval parameter represents the frequency at which the IoT devices send their data to the fog nodes or cloud servers. As the snapshot interval increases, the frequency of
data transmission decreases. With a snapshot interval of 1, the IoT devices are sending data continuously. As the number of devices increases, the energy consumption of both with fog and without fog scenarios increases slightly, but the with fog mean remains consistently higher than the without fog mean. With a snapshot interval of 5, the IoT devices are sending data less frequently, which results in reduced energy consumption. In this case, the energy consumption of the with fog scenario is consistently lower than the without fog scenario, which demonstrates the energy efficiency advantages of using fog computing. With a snapshot interval of 10, the IoT devices send data even less frequently, and the difference in energy consumption between the with fog and without fog scenarios becomes more pronounced. This result further emphasizes the benefits of using fog computing in terms of energy efficiency.

**Number of Devices**

The number of devices parameter refers to the number of telehealth IoT devices in the network. As the number of devices increases, the energy consumption for both with fog and without fog scenarios tends to increase as well. This is expected, as more devices lead to higher data transmission and processing loads. However, the increase in energy consumption is consistently smaller in the with fog scenario compared to the without fog scenario across all snapshot intervals. This shows that the proposed fog-based model is more scalable and can better handle the energy requirements of a growing number of devices.

**With Fog Mean and Without Fog Mean**

The with fog mean and without fog mean parameters represent the average energy consumption in the scenarios with and without fog computing, respectively. Across all snapshot intervals and several devices, the with fog mean is generally lower than the without fog mean, indicating that the fog-based model is more energy-efficient than the cloud-only model.

**With Fog SD and Without Fog SD**

The with fog SD and without fog SD parameters represent the SD of the energy consumption in the scenarios with and without fog computing, respectively. In general, the SD values are lower in the with fog scenario compared to the without fog scenario. This suggests that the energy consumption is more consistent and less variable in the fog-based model, which could lead to more predictable and stable system performance.

**With Fog 95% CI and Without Fog 95% CI**

The CI in the simulation code is a range within which a certain percentage of the population parameter is expected to lie, with a specified level of confidence. In the context of the provided simulation results, the 95% CIs represent the range within which the true mean performance of the system (either with or without fog computing) is likely to fall, with a certain level of confidence, typically 95%.

A 95% CI is calculated using the sample mean, sample SD, and sample size. The formula for a 95% CI is:

\[
CI = \text{sample mean} \pm (1.96 \times \frac{\text{sample SD}}{\sqrt{\text{sample size}}})
\]

The 95% CI helps to quantify the uncertainty in the estimation of the true mean performance. A narrower 95% CI indicates a more precise estimate, while a wider interval suggests more uncertainty.

**Analysis of Results**

Table 1 contains the summary of statistical results.

<table>
<thead>
<tr>
<th>Snapshot interval</th>
<th>Number of devices</th>
<th>With fog, mean (SD)</th>
<th>With fog, 95% CI</th>
<th>With fog, mean (SD)</th>
<th>Without fog, mean (SD)</th>
<th>Without fog, 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10</td>
<td>90.43 (0.45)</td>
<td>90.11-90.76</td>
<td>89.74 (0.05)</td>
<td>89.69-89.79</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>20</td>
<td>90.53 (0.33)</td>
<td>90.30-90.77</td>
<td>89.74 (0.06)</td>
<td>89.69-89.79</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>30</td>
<td>90.61 (0.23)</td>
<td>90.45-90.78</td>
<td>89.74 (0.04)</td>
<td>89.71-89.78</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>40</td>
<td>90.55 (0.24)</td>
<td>90.38-90.72</td>
<td>89.76 (0.05)</td>
<td>89.71-89.90</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>10</td>
<td>87.39 (0.70)</td>
<td>86.89-87.89</td>
<td>86.04 (0.13)</td>
<td>85.94-86.13</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>20</td>
<td>86.59 (0.21)</td>
<td>86.44-86.73</td>
<td>85.91 (0.06)</td>
<td>85.86-85.95</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>30</td>
<td>87.02 (0.46)</td>
<td>86.70-87.34</td>
<td>86.01 (0.09)</td>
<td>85.95-86.08</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>40</td>
<td>87.30 (0.27)</td>
<td>87.12-87.50</td>
<td>86.00 (0.07)</td>
<td>85.95-86.05</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>10</td>
<td>82.85 (0.73)</td>
<td>82.34-83.38</td>
<td>81.36 (0.11)</td>
<td>81.27-81.44</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>20</td>
<td>83.28 (0.63)</td>
<td>82.83-83.72</td>
<td>81.42 (0.11)</td>
<td>81.33-81.50</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>30</td>
<td>82.62 (0.59)</td>
<td>82.80-83.03</td>
<td>81.33 (0.12)</td>
<td>81.24-81.43</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>40</td>
<td>82.7 (0.37)</td>
<td>82.43-82.95</td>
<td>81.36 (0.07)</td>
<td>81.31-81.41</td>
<td></td>
</tr>
</tbody>
</table>

Here is a step-by-step analysis of the results (Table 1):

1. Observe the “With fog, mean (SD)” and “Without fog, mean (SD)” columns for each combination of “Snapshot interval” and “Number of devices.” In all cases, the with fog mean is higher than the without fog mean, indicating that, on average, the remaining energy is higher when using fog computing.

2. Look at the 95% CIs for both “with fog” and “without fog” scenarios. If the 95% CIs do not overlap, it suggests that the difference in energy remaining between the 2 scenarios is statistically significant. For example, in the first row
(snapshot interval: 1, number of devices: 10), the “with fog, 95% CI” is 87.98-89.45, and the “without fog, 95% CI” is 84.90-87.47. Since these intervals do not overlap, there is strong evidence that using fog computing leads to significantly higher energy remaining for this specific combination of parameters.

3. Compare the width of the 95% CIs for each scenario. A narrower 95% CI indicates a more precise estimate of the true population mean. For most 95% CI values, the “with fog, 95% CI” is narrower than the “without fog, 95% CI,” suggesting that the “with fog” scenario has a more precise estimate.

4. Analyze the trends as the number of devices increases within each snapshot interval. In general, the energy remaining in both scenarios decreases as the number of devices increases. However, the rate of decrease seems to be lower when using fog computing.

5. Observe the trends as the snapshot interval increases for each group of devices. As the snapshot interval increases, the energy remaining for both scenarios decreases, suggesting that less frequent snapshots may lead to less energy conservation. However, the “with fog” scenario consistently results in higher energy remaining compared to the “without fog” scenario, regardless of the snapshot interval.

In conclusion, based on the analysis of the means and 95% CIs, it appears that using fog computing is beneficial for conserving energy, especially when the number of devices and the snapshot intervals increase. The difference in energy remaining is statistically significant in most cases, and the “with fog” scenario consistently outperforms the “without fog” scenario.

Table 2. Sensitivity analysis with energy cost.

<table>
<thead>
<tr>
<th>Energy cost</th>
<th>With fog, mean (SD)</th>
<th>Without fog, mean (SD)</th>
<th>Mean difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.20</td>
<td>94 (5)</td>
<td>91 (4)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.26</td>
<td>93 (6)</td>
<td>92 (5)</td>
<td>0.33</td>
</tr>
<tr>
<td>0.32</td>
<td>94 (4)</td>
<td>92 (7)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.38</td>
<td>93 (5)</td>
<td>91 (2)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.44</td>
<td>94 (3)</td>
<td>91 (5)</td>
<td>2.75</td>
</tr>
<tr>
<td>0.5</td>
<td>93 (2)</td>
<td>91 (4)</td>
<td>1.71</td>
</tr>
<tr>
<td>0.56</td>
<td>95 (2)</td>
<td>92 (2)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.62</td>
<td>94 (3)</td>
<td>92 (4)</td>
<td>1.37</td>
</tr>
<tr>
<td>0.68</td>
<td>93 (2)</td>
<td>92 (2)</td>
<td>0.68</td>
</tr>
<tr>
<td>0.74</td>
<td>94 (4)</td>
<td>92 (3)</td>
<td>1.02</td>
</tr>
<tr>
<td>0.80</td>
<td>95 (6)</td>
<td>92 (3)</td>
<td>2.06</td>
</tr>
</tbody>
</table>

Based on the sensitivity analysis of energy cost, the mean energy remaining for IoT devices with fog nodes is consistently higher than that of devices without fog nodes across all energy cost values. This indicates that IoT devices with fog nodes perform better in terms of energy consumption as compared to devices without fog nodes.

Table 3 compares the mean energy remaining for IoT devices with and without fog nodes for each latency parameter value. The “Mean difference” column shows the difference in mean energy remaining, with positive values indicating that devices with fog nodes have higher energy remaining compared to those without fog nodes. In the with fog scenario, the mean energy remaining for devices with fog nodes stays relatively stable, ranging from a minimum of 93 to a maximum of 95 across different energy costs. In the without fog scenario, the mean energy remaining for devices without fog nodes also remains relatively stable, ranging from a minimum of 91 to a maximum of 92 across different energy costs.

Table 3. Sensitivity analysis with latency parameter.

<table>
<thead>
<tr>
<th>Latency parameter</th>
<th>With fog, mean (SD)</th>
<th>Without fog, mean (SD)</th>
<th>Mean difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>94 (5)</td>
<td>91 (4)</td>
<td>1.72</td>
</tr>
<tr>
<td>2</td>
<td>93 (6)</td>
<td>92 (5)</td>
<td>0.33</td>
</tr>
<tr>
<td>3</td>
<td>94 (4)</td>
<td>92 (7)</td>
<td>1.72</td>
</tr>
<tr>
<td>4</td>
<td>93 (5)</td>
<td>91 (2)</td>
<td>1.72</td>
</tr>
<tr>
<td>5</td>
<td>94 (3)</td>
<td>91 (5)</td>
<td>2.75</td>
</tr>
<tr>
<td>6</td>
<td>93 (2)</td>
<td>91 (4)</td>
<td>1.71</td>
</tr>
<tr>
<td>7</td>
<td>95 (2)</td>
<td>92 (2)</td>
<td>1.72</td>
</tr>
<tr>
<td>8</td>
<td>94 (3)</td>
<td>92 (4)</td>
<td>1.37</td>
</tr>
<tr>
<td>9</td>
<td>93 (2)</td>
<td>92 (2)</td>
<td>0.68</td>
</tr>
<tr>
<td>10</td>
<td>94 (4)</td>
<td>92 (3)</td>
<td>1.02</td>
</tr>
<tr>
<td>11</td>
<td>95 (6)</td>
<td>92 (3)</td>
<td>2.06</td>
</tr>
</tbody>
</table>

Based on the sensitivity analysis of latency parameter, the mean energy remaining for IoT devices with fog nodes is consistently higher than that of devices without fog nodes across all latency values. This indicates that IoT devices with fog nodes perform better in terms of energy consumption as compared to devices without fog nodes.
relatively stable, ranging from a minimum of 91 to a maximum of 93 across different latency values.

Based on the sensitivity analysis of latency, the mean energy remaining for IoT devices with fog nodes is consistently higher than that of devices without fog nodes across all latency parameter values. This indicates that IoT devices with fog nodes perform better in terms of energy consumption as compared to devices without fog nodes.

Table 4 compares the mean energy remaining for IoT devices with and without fog nodes for each idle power value. The “Mean difference” column shows the difference in mean energy remaining, with positive values indicating that devices with fog nodes have higher energy remaining compared to those without fog nodes. In the with fog scenario, the mean energy remaining for devices with fog nodes stays relatively stable, ranging from a minimum of 93 to a maximum of 95 across different idle power values. In the without fog scenario, the mean energy remaining for devices without fog nodes also remains relatively stable, ranging from a minimum of 90 to a maximum of 92 across different idle power values.

Table 3. Sensitivity analysis with latency.

<table>
<thead>
<tr>
<th>Latency parameter</th>
<th>With fog, mean (SD)</th>
<th>Without fog, mean (SD)</th>
<th>Mean difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.20</td>
<td>94 (4)</td>
<td>92 (4)</td>
<td>2.06</td>
</tr>
<tr>
<td>0.26</td>
<td>93 (3)</td>
<td>91 (4)</td>
<td>2.06</td>
</tr>
<tr>
<td>0.32</td>
<td>94 (2)</td>
<td>93 (1)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.38</td>
<td>95 (2)</td>
<td>92 (2)</td>
<td>2.06</td>
</tr>
<tr>
<td>0.44</td>
<td>94 (2)</td>
<td>92 (2)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.5</td>
<td>94 (3)</td>
<td>91 (3)</td>
<td>2.4</td>
</tr>
<tr>
<td>0.56</td>
<td>94 (4)</td>
<td>92 (5)</td>
<td>2.06</td>
</tr>
<tr>
<td>0.62</td>
<td>94 (2)</td>
<td>92 (2)</td>
<td>2.06</td>
</tr>
<tr>
<td>0.68</td>
<td>94 (3)</td>
<td>92 (3)</td>
<td>1.71</td>
</tr>
<tr>
<td>0.74</td>
<td>94 (5)</td>
<td>93 (3)</td>
<td>0.68</td>
</tr>
<tr>
<td>0.80</td>
<td>94 (2)</td>
<td>92 (4)</td>
<td>0.68</td>
</tr>
</tbody>
</table>

Table 4. Sensitivity analysis with idle power.

<table>
<thead>
<tr>
<th>Idle power</th>
<th>With fog, mean (SD)</th>
<th>Without fog, mean (SD)</th>
<th>Mean difference</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>95 (2)</td>
<td>92 (2)</td>
<td>2.06</td>
</tr>
<tr>
<td>0.6</td>
<td>94 (2)</td>
<td>92 (4)</td>
<td>1.71</td>
</tr>
<tr>
<td>0.7</td>
<td>95 (3)</td>
<td>92 (4)</td>
<td>2.41</td>
</tr>
<tr>
<td>0.8</td>
<td>93 (2)</td>
<td>90 (3)</td>
<td>1.72</td>
</tr>
<tr>
<td>0.9</td>
<td>94 (4)</td>
<td>92 (4)</td>
<td>1.03</td>
</tr>
<tr>
<td>1.0</td>
<td>93 (5)</td>
<td>91 (4)</td>
<td>1.02</td>
</tr>
<tr>
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<td>95 (4)</td>
<td>92 (6)</td>
<td>1.71</td>
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<tr>
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<td>94 (4)</td>
<td>91 (5)</td>
<td>2.06</td>
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<td>95 (2)</td>
<td>91 (2)</td>
<td>2.75</td>
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<tr>
<td>1.5</td>
<td>94 (2)</td>
<td>92 (4)</td>
<td>1.72</td>
</tr>
</tbody>
</table>

Based on the sensitivity analysis of idle power, the mean energy remaining for IoT devices with fog nodes is consistently higher than that of devices without fog nodes across all idle power values. This indicates that IoT devices with fog nodes perform better in terms of energy consumption as compared to devices without fog nodes.

Table 5 compares the mean energy remaining for IoT devices with and without fog nodes for each transmit power value. The “Mean difference” column shows the difference in mean energy remaining, with positive values indicating that devices with fog nodes have higher energy remaining compared to those without fog nodes. In the with fog scenario, the mean energy remaining for devices with fog nodes stays relatively stable, ranging from a minimum of 94 to a maximum of 96 across different transmit power values. In the without fog scenario, the mean energy remaining for devices without fog nodes also remains relatively stable, ranging from a minimum of 91 to a maximum of 92 across different transmit power values.
Table 5. Sensitivity analysis with transmit power.

<table>
<thead>
<tr>
<th>Transmit power</th>
<th>With fog, mean (SD)</th>
<th>Without fog, mean (SD)</th>
<th>Mean difference</th>
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<tr>
<td>0.5</td>
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<tr>
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<td>94 (2)</td>
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<td>1.5</td>
<td>95 (2)</td>
<td>91 (1)</td>
<td>3.44</td>
</tr>
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</table>

Based on the sensitivity analysis of transmit power, the mean energy remaining for IoT devices with fog nodes is consistently higher than that of devices without fog nodes across all transmit power values. This indicates that IoT devices with fog nodes perform better in terms of energy consumption as compared to devices without fog nodes.

Ethical Considerations

The study did not apply for any ethical approval, as the research did not involve any human participants or animals [44].

Discussion

Overview

The simulation study results indicate that the proposed energy-saving model could be effective in reducing energy consumption in real-world telehealth scenarios. Key findings include the following:

1. Scalability: the model demonstrates the ability to accommodate an increasing number of IoT devices without compromising performance, energy efficiency, or quality of health care services.

2. Task allocation algorithm: the proposed task allocation algorithm outperforms other algorithms in terms of energy efficiency and data processing efficiency, indicating its effectiveness in balancing the workload between fog nodes and cloud servers.

3. Energy consumption metrics: the overall energy consumption is reduced across all levels, demonstrating the success of the model’s energy-saving strategies, such as adaptive power management, data compression, and network optimization.

The code and methodology described aim to simulate an IoT network with different components (IoT devices, fog nodes, and cloud servers) and analyze the impact of fog nodes on energy consumption. The code creates and connects these components and simulates data transmission, storage, and energy consumption for IoT devices, fog nodes, and cloud servers. The simulation results are analyzed to understand the network behavior and demonstrate the potential benefits of using fog nodes for energy efficiency.

Our novel energy-efficient model integrates fog and cloud computing paradigms to optimize data processing for telehealth IoT devices without compromising real-time health care services. This stands out from previous works by enabling localized data processing through the incorporation of fog computing. This intermediary layer, situated between IoT devices and cloud servers, effectively reduces latency and data transfer overhead. The concurrent use of public and private cloud computing further fortifies the system’s infrastructure, allowing for the handling of large data volumes and resource-intensive computations. The model enables localized data processing by incorporating fog computing as an intermediary layer between IoT devices and public or private cloud servers, effectively reducing latency and data transfer overhead. Simultaneously, public and private cloud computing provides a robust infrastructure for handling large data volumes and performing resource-intensive computations. The primary goal of this model is to minimize energy consumption through intelligent task allocation between fog nodes and cloud servers, by considering their computational capacity and proximity to IoT devices. This task allocation process also considers various sensitivity and priority levels within the health care context, ensuring prompt responses to critical and high-sensitivity requests. Our innovative model strategically integrates fog and cloud computing, aiming to establish an energy-efficient telehealth IoT system capable of adeptly managing data processing and delivering real-time health care services, accommodating various levels of sensitivity and priorities. While these aspirations suggest promising opportunities for further optimization and diverse applications within health care contexts, it is crucial to note that the subsequent simulation method serves to objectively assess the model’s effectiveness and efficiency. The empirical evidence derived from the simulation provides a foundation for a more nuanced understanding of the model’s capabilities and potential benefits. This is because exploring diverse large-scale network topologies is rarely feasible in the real world. Although the requirements for such a simulator are straightforward—providing a detailed,
accurate, and granular model of all components—implementing corresponding simulators demands considerable effort.

The primary strength of our model lies in its holistic approach toward minimizing energy consumption. The intelligent task allocation mechanism, considering computational capacity and proximity to IoT devices, ensures a fine balance. Furthermore, the incorporation of sensitivity and priority levels within the health care context enhances the model’s responsiveness to critical requests. The synergistic integration of fog and cloud computing contributes to the creation of an energy-efficient telehealth IoT system capable of real-time data processing in accordance with varying sensitivity levels and priorities.

Despite the positive outcomes, several limitations should be acknowledged. (1) Simulation environment realism: the simulation, while essential for its controlled environment, may not perfectly mirror real-world complexities. Variations in network behaviors and external factors may influence results differently in practical implementations. (2) Sensitivity analysis scope: the sensitivity analysis, while comprehensive, focused on specific parameters such as energy cost, latency, idle power, and transmit power. Additional parameters and their potential interactions may provide a more nuanced understanding of the model’s behavior. (3) Simplifications in simulation: certain simplifications, inherent in simulation models, may oversimplify the intricacies of a live telehealth IoT deployment. Real-world complexities such as device failures, communication errors, or dynamic changes in the environment are challenging to fully capture.

To address these limitations and advance the research, the following suggestions should be considered. (1) Future studies should aim for more realistic simulation environments, incorporating dynamic factors and diverse network topologies to enhance the model’s external validity. (2) Expanding the scope of sensitivity analysis to include a broader range of parameters and exploring their interactions could provide a more comprehensive understanding of the model’s performance under diverse conditions. (3) The development of more sophisticated simulators, despite their challenges, remains crucial. Detailed, accurate, and granular models of all components can better simulate the intricacies of large-scale IoT-fog-cloud systems.

While our model exhibits significant promise in reducing energy consumption and enhancing data processing efficiency in telehealth IoT scenarios, ongoing refinement and exploration of diverse scenarios will contribute to its continued evolution and real-world applicability.

Conclusion
This paper provides a compelling model for the use of fog and cloud computing–based platforms in telehealth IoT deployments to reduce energy consumption, improve data processing efficiency, and maintain high-quality health care services. The model leverages the strengths of both fog and cloud computing paradigms to address the challenges associated with large-scale telehealth IoT deployments, such as energy consumption, data processing efficiency, latency, security, and privacy. The simulation results show that the proposed fog-based model significantly reduces energy consumption compared to the cloud-only model while maintaining high-quality data processing and transmission. Moreover, the methodology described in this paper provides a comprehensive approach to analyzing network performance and energy consumption, which includes examining the impact of various parameters, such as the number of devices, fog node deployment, task allocation algorithm, energy consumption metrics, and performance metrics. Sensitivity analyses were conducted with respect to energy cost, latency, idle power, and transmit power, consistently showing that IoT devices with fog nodes had higher mean energy remaining compared to devices without fog nodes. This approach allows for a more detailed understanding of the network behavior and potential bottlenecks and provides insights into how to optimize the model to be more resilient and efficient.

The simulation results and methodology demonstrate the effectiveness of the proposed model and provide a roadmap for future research in this area. We demonstrated the effectiveness of the proposed model in reducing energy consumption while, more importantly, ensuring efficient data processing and maintaining the quality of health care services. The proposed model can help health care providers and stakeholders improve patient care and outcomes while reducing costs and energy consumption.

Acknowledgments
This work was supported by the National Natural Science Foundation of Canada and the Department of Computer Science, University of Victoria.

Conflicts of Interest
None declared.

References


34. Aazam M, Huh EN. Fog computing and smart gateway based communication for cloud of things. 2015 Presented at: 2015 International Conference on Future Internet of Things and Cloud (FiCloud); August 24-26, 2015; Rome, Italy p. 464-470.

Abbreviations

IoT: Internet of Things  
YAFS: Yet Another Fog Simulator
information, a link to the original publication on https://biomedeng.jmir.org/, as well as this copyright and license information must be included.
An Engineering Alternative to Lockdown During COVID-19 and Other Airborne Infectious Disease Pandemics: Feasibility Study

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Abstract

Background: Now and in the future, airborne diseases such as COVID-19 could become uncontrollable and lead the world into lockdowns. Finding alternatives to lockdowns, which limit individual freedoms and cause enormous economic losses, is critical.

Objective: The purpose of this study was to assess the feasibility of achieving a society or a nation that does not require lockdown during a pandemic due to airborne infectious diseases through the mass production and distribution of high-performance, low-cost, and comfortable powered air purifying respirators (PAPRs).

Methods: The feasibility of a social system using PAPR as an alternative to lockdown was examined from the following perspectives: first, what PAPRs can do as an alternative to lockdown; second, how to operate a social system utilizing PAPR; third, directions of improvement of PAPR as an alternative to lockdown; and finally, balancing between efficiency of infection control and personal freedom through the use of Internet of Things (IoT).

Results: PAPR was shown to be a possible alternative to lockdown through the reduction of airborne and droplet transmissions and through a temporary reduction of infection probability per contact. A social system in which individual constraints imposed by lockdown are replaced by PAPRs was proposed, and an example of its operation is presented in this paper. For example, the government determines the type and intensity of the lockdown and activates it. At that time, the government will also indicate how PAPR can be substituted for the different activity and movement restrictions imposed during a lockdown, for example, a curfew order may be replaced with the permission to go outside if wearing a PAPR. The following 7 points were raised as directions for improvement of PAPR as an alternative method to lockdown: flow optimization, precise differential pressure control, design improvement, maintenance method, variation development such as booth type, information terminal function, and performance evaluation method. In order to achieve the effectiveness and efficiency in controlling the spread of infection and the individual freedom at a high level in a social system that uses PAPRs as an alternative to lockdown, it was considered effective to develop a PAPR wearing rate network management system utilizing IoT.

Conclusions: This study shows that using PAPR with infection control ability and with less economic and social damage as an alternative to nationwide lockdown is possible during a pandemic due to airborne infectious diseases. Further, the efficiency of the government’s infection control and each citizen’s freedom can be balanced by using the PAPR wearing rate network management system utilizing an IoT system.

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KEYWORDS
COVID-19; airborne infectious diseases; lockdown; powered air purifying respirator (PAPR); infectious dose; airborne transmission; emergency evacuation; herd immunity; aerosol; air; quality; infection control; infectious; respiratory; purifier; purifiers; purifying; respirator; respirators; device; devices; airborne
Introduction

For more than 3 years, herd immunity has been pursued worldwide through vaccination as a countermeasure against COVID-19, in addition to new lifestyle measures (social distancing, wearing of face masks, washing of hands, etc) [1,2]. However, due to the emergence of new variants, the spread of infection has sometimes been uncontrollable in different parts of the world. Each time, lockdown has been implemented to temporarily buy time, causing great economic loss and restriction of freedom for individuals, businesses, and society [3-7]. In this study, lockdown is defined as restricting the actions and activities of people and businesses to temporarily slow the spread of infection and buy time for other measures such as herd immunity through vaccination. A strong lockdown includes orders with criminal penalties that prohibit going out, working, and doing business, and a weak lockdown includes voluntary requests to refrain from going out, working, and doing business. The type and strength of lockdowns are determined by the government on a case-by-case basis, considering the impact on infection control and harm to society, depending on the situation of the outbreak of infection at the time [8,9].

Even in the current situation, the rapid development and delivery of effective vaccines against new variants of the coronavirus (SARS-CoV-2) and new airborne viruses that may emerge in succession is not well assured. In the future, it is likely that we will continue to be in a situation where we do not know when a lockdown will again be necessary around the world [10,11]. In light of this, alternatives to lockdown that can cause less economic damage, avoid restrictions on freedom of action, and other disadvantages to individuals, companies, and societies would be beneficial. Among the modes of COVID-19 transmission, contact and oral infections are relatively easy to prevent by environmental hygiene, including handwashing and food hygiene management. Droplet infection (particle size ≥100 µm) is thought to be preventable by social distancing (droplets fall by gravity) and by wearing a mask. Currently, airborne transmission via aerosols (particle size <100 µm) is thought to be the main route of infection [12,13]. Although the infectious dose of COVID-19, that is, the number of ingested viruses required for infection, is not well known [14], it is estimated to be in the range of 300 to 2000 virions [15]. It is believed that the possibility of viral infection can be effectively reduced by shielding aerosols that may contain viruses. This paper discusses alternative means to lockdown, assuming that contact and oral infections are prevented by environmental hygiene and food hygiene management and that only airborne and droplet infections remain as infection routes.

Methods

The feasibility of a social system utilizing powered air purifying respirators (PAPRs) as an alternative to lockdown was examined from the following perspectives: (1) what PAPR can do as an alternative to lockdown, (2) how to operate a social system utilizing PAPR, (3) direction of improvement of PAPR as an alternative to lockdown, and (4) balance between efficiency of infection control and personal freedom through the use of Internet of Things (IoT).

What PAPR Can Do as an Alternative to Lockdown

The PAPR for practical medical use is a device that drastically reduces the number of viruses inhaled by the wearer and effectively reduces the risk of infection. Medical PAPRs are used by medical personnel working in high-risk environments [16,17]. The assigned protection factor (APF), as defined by the National Institute for Occupational Safety and Health in the United States, is widely used as an indicator of the shielding performance of respiratory protection devices, including PAPRs [18]. APF is defined as the external concentration/internal concentration of the target particles (aerosols). For a medical face mask, APF=10 is given when a person who has been trained to wear it wears it completely with no gaps between the mask and the facial surface. However, 3M PAPR is rated at APF=1000 and is considered to have excellent protective performance [19]. In other words, the concentration of virus-containing aerosols can be reduced to 1/10 or less with a full-face mask when worn without gaps, while it is reduced to 1/1000 or less with 3M PAPR. Therefore, high-performance PAPRs can be used as an alternative to the movement restrictions and activity restrictions imposed by lockdowns used as a countermeasure against COVID-19 and other airborne infectious diseases in future pandemics. The necessary conditions for an alternative to lockdown are that it should have the same deterrent effect on the spread of infection as lockdown, and the economic damage, activity restrictions, and other disadvantages to individuals and society should be smaller compared to those during lockdown.

The effective reproduction number \( R_e \) is the average number of secondary cases per infectious case in a population of both susceptible and nonsusceptible individuals. In this population, \( R_e<1 \) means converging, \( R_e=1 \) means stationary, and \( R_e>1 \) means expanding [20]. The effective reproduction number \( R_e \) can be expressed schematically by the following equation:

\[
R_e = \beta \times k \times D
\]

where \( \beta \) is the probability of infection being transmitted during a contact, \( k \) (contact/day)=contact rate in the host population, and \( D \) (day)=duration of infectiousness.

To control the infection of the whole society, it is sufficient to set the effective reproduction number \( R_e \) to <1. To do this, we should reduce the above \( \beta, k, \) and \( D \). Vaccines are expected to reduce \( \beta \) in the long term. Lockdown is expected to play a role in temporarily reducing \( k \). In this study, PAPR is expected to play a role in temporarily reducing \( \beta \) by reducing the airborne and droplet transmission. During COVID-19, the maximum \( R_e \) value reported worldwide was around 5 [21,22]. If lockdown is used as a means of correcting \( R_e=5 \) to \( R_e=1 \), it is sufficient if the lockdown makes \( k \) 1/5 of its current value. Alternatively, if PAPRs are used as an alternative to lockdown, it would be sufficient if citizens could wear PAPRs to reduce \( \beta \) to 1/5 of its current value. It is worth noting here that the degree of reduction in \( k \) and \( \beta \) may be about 1/5 rather than 1/100 or 1/1000. The performance of the existing PAPRs was evaluated. Figure 1 shows the specifications of 3 existing PAPRs.
Currently Existing PAPRs

**Medical PAPR: Versaflo TR-301N+**

This PAPR is marketed as a medical PAPR. The waist unit (model TR-301N+; 3M Corp) contains a nonwoven fabric filter (model TR-3712N), pump, and battery (model TR-332). Purified air is pumped from the waist unit into the hood (model S-133L) through a flexible hose (model PSD-0225). The outline of this PAPR is as follows. Only air purified by a high-performance nonwoven filter is introduced into the hood by a pump. Since positive pressure is naturally maintained inside the hood, outside air is prevented from entering even if there is a gap between the hood seal and the face and the head. Thus, the air, which the wearer breathes, is only the air purified by the nonwoven filter. As for the face mask, since the inside of the mask becomes negative pressure during inhalation, if there is a gap between the face and the mask, the outside air will enter directly through the gap during the wearer’s inhalation [23,24]. Therefore, PAPR is structurally capable of shielding most of the aerosols present in the ambient air. This is supported by the fact that this PAPR has a high value of APF=1000. The flow rate can be selected in 2 stages, that is, high and low, and the specified flow rates are approximately 6.5 cubic feet per minute (180 L/min) and 7.2 cubic feet per minute (200 L/min), respectively [25].

**PAPR Prototype With a Simple Structure**

This PAPR is a prototype developed as a low-cost PAPR that has a simple structure similar to the abovementioned commercially available medical PAPR. The specifications of the air purification characteristics in this prototype are as follows [26].

1. Only air purified by a high-efficiency nonwoven filter is pumped into the hood.

2. Because positive pressure is naturally maintained inside the hood, outside air is prevented from entering even if there is a gap between the hood seal and the face.

3. The exhaust is natural exhaust from a thin nonwoven fabric filter due to the positive pressure inside the hood. Even if the wearer is infected and emits droplets or aerosols containing the virus, it is possible to prevent some of the external emissions. The first 2 characteristics mentioned above are the same as those for the medical PAPR mentioned above (TR-301N+). In the air supply side, a nonwoven fabric filter—high-efficiency particulate air filter—which can filter 99.97% or more of aerosols down to 0.3 \(\mu\)m or larger is used on the air supply side. For aerosols containing viruses, it is considered sufficient to target aerosols with a particle size of 0.3 \(\mu\)m or larger [27,28].

A high-performance PAPR can be assembled at a total parts cost of approximately US $40 with a simple configuration of only a high performance filter, battery, and pump.

**PAPR Prototype With a Controller**

This PAPR is a prototype developed as a high-performance PAPR equipped with a controller/computer for measurement and control [29]. The specifications for the air purification characteristics are the same as those in the PAPR prototype with a simple structure. In addition to the characteristics of the PAPR prototype with a simple structure, a controller (on-board computer) and sensors (2 CO₂ concentration sensors and 1 differential pressure sensor) are added, and the pump is controlled by means of the pulse width modulation control. In this PAPR, the pump output is adjusted and controlled according to the output of the differential pressure sensor so as to suppress the internal pressure fluctuations due to breathing, that is, higher pressure during exhalation and lower pressure during inspiration. Operating parameters can be set and monitored using a
smartphone. It is also possible to connect to the internet via the smartphone. As a result, it is possible to connect to the PAPR wearing rate network management system. In addition, by installing a pump and filter on the exhaust side and making it a differential type, it is possible to set the internal pressure to either positive pressure or negative pressure. Thus, it is possible to set the positive pressure setting to protect the wearer from the outside and to set the negative pressure setting to protect the outside from the wearer. It is possible to manufacture this PAPR with a total parts cost of approximately US $300; this PAPR allows pump control based on sensor signals and settings and monitoring of the operating parameters by using a smartphone.

How to Operate a Social System Utilizing PAPR

A case in which a fixed percentage of the population wears PAPRs is considered. Quantitative evaluation of the reduction in the aforementioned $(\beta)$ (infection probability per contact) by wearing PAPR and quantitative evaluation of the reduction in $(R_s)$ by wearing PAPR are considered to require empirical and social experiments, as described below, because they involve humans. However, as shown below, quantitative evaluation is possible under limited conditions. Assuming that the aerosols and droplet shielding rate by PAPRs is the same for all particle sizes of aerosols and droplets, the shielding rate can be expressed as follows: (1) shielding rate for aerosols and droplets in the air supply $(R_{S,in})$ and (2) shielding rate for aerosols and droplets in the exhaust air $(R_{S,ex})$. Assume that the reduction rate of the probability of the wearer himself/herself becoming infected and the reduction rate of the probability of the wearer infecting others by wearing PAPR are as follows: (3) reduction rate of the probability that the wearer will be infected by aerosols and droplets in the air supply $(R_{I,in})$ and (4) reduction rate of the probability of infecting others by aerosols and droplets in exhaust air $(R_{I,ex})$. It is difficult to quantitatively determine the relationship between (1) and (3) and between (2) and (4) above. However, the following limited arrangement can be made. For the air supply side, the following can be said.

1. If the aerosol and droplet shielding ratio of PAPR is perfect $(R_{S,in}=1)$, wearing PAPR will reduce the probability of infection by means of aerosols and droplets in the supply air by 100% $(R_{I,in}=1)$.
2. If the aerosol and droplet shielding ratio of PAPR is nothing $(R_{S,in}=0)$, wearing PAPR will reduce the probability of infection by means of aerosols and droplets in the supply air by nothing $(R_{I,in}=0)$.
3. In the intervals of $0<R_{S,in}<1$ and $0<R_{I,in}<1$, there is a positive correlation between $R_{S,in}$ and $R_{I,in}$.

For the exhaust side, the following can be said.

4. If the aerosol and droplet shielding ratio of PAPR is perfect $(R_{S,ex}=1)$, wearing PAPR will reduce the probability of infecting others by means of the aerosols and droplets in the exhaust air by 100% $(R_{I,ex}=1)$.
5. If the aerosol and droplet shielding ratio of PAPR is nothing $(R_{S,ex}=0)$, wearing PAPR will reduce the probability of infecting others by means of the aerosols and droplets in the exhaust air by nothing $(R_{I,ex}=0)$.
6. In the intervals of $0<R_{S,ex}<1$ and $0<R_{I,ex}<1$, there is a positive correlation between $R_{S,ex}$ and $R_{I,ex}$.

A social group was assumed to be completely free from contact and oral infections, and airborne and droplet infections were the only routes of infection. It is assumed that a certain percentage of the population in the social group always wears a PAPR. The performance of PAPR is assumed to be as follows.

1. The shielding rate of aerosols and droplets in the air supply side is 100% $(R_{S,in}=1)$. As a result, the reduction rate of the probability of infection by aerosol and droplets in the air supply is 100% $(R_{I,in}=1)$.
2. The shielding rate of aerosols and droplets in the exhaust side is the same as that of the face mask used in the population at that time.
3. When the above PAPR is worn by a percentage of people in $W_R$, the following relationship is established between the effective reproduction number $R_t$ immediately before the start of wearing and the modified effective reproduction number $R_{\text{im}}$ immediately after the start of wearing.

$$W_R = 1 - R_{\text{im}} / R_t$$

*Figure 2* shows the relationship between the effective reproduction number $R_t$ at the time in question and the required wearing rate $W_{R,\text{required}}$, which is required to achieve the target effective reproduction number $R_{\text{im}}$ of 0.5, 0.9, and 1.0. For example, consider an event in which a certain percentage of the population is wearing PAPR at all the time. As an example of a situation of severe infection spread, consider the case where $R_t=2$ immediately before the start of the event. In this case, to achieve the target effective reproduction number $R_{\text{im}}$ of 1.0, 0.9, and 0.5, 50%, 55%, and 75% of the population should wear PAPRs at all times, respectively. The above simulation targets a social group in which airborne and droplet infections are the only routes of infection and an extreme setting in which PAPR is worn at all times. Future studies should consider more realistic settings that suit the conditions of daily life. For example, the situations in which the effects of not wearing PAPR should be considered, including contact with family members in the home, eating, drinking, washing, and bathing.
Direction of Improvement of PAPR as an Alternative to Lockdown

As the directions of improvements of PAPR as an alternative to lockdown, the following 7 points are proposed and discussed: (1) flow path optimization, (2) precise pressure control by fluid modeling, (3) improved design, (4) maintenance method, (5) variations suitable for different places of use and activity contents, (6) PAPR with information terminal function, and (7) evaluation indicators and evaluation methods.

Balance Between Efficiency of Infection Control and Personal Freedom Through the Use of IoT

In order to achieve both (1) effectiveness and efficiency in controlling the spread of infection and (2) individual freedom (limiting the obligation to wear PAPRs to the minimum necessary) at a high level in a social system that uses PAPRs as an alternative to lockdown, it is considered effective to develop a PAPR wearing rate network management system as shown in Figure 3.
1. PAPR (helmet type, booth type, etc) is connected to the wearer’s smartphone via Bluetooth.

2. The smartphone is connected to the internet and connected to the PAPR wearing rate network management system server operated by the government.

3. The government will be able to monitor, record, and manage each citizen’s PAPR wearing rate along with smartphone location information by using the system.

Wearers (citizens) can display the electronic proof of their wearing rate on their smartphones provided by the system. Various parameters can be considered for the PAPR wearing rate ($W_R$). As simple examples, the following definitions of wearing rate ($W_R$) can be considered.

$$W_R = \frac{\text{time spent outside with PAPR}}{\text{time spent outside}}$$

Instead of the PAPR wearing rate $W_R$, time spent outside without PAPR $T_{WT}$ can be considered.

$$T_{WT} = \text{time spent outside without PAPR}$$

Instead of the PAPR wearing rate $W_R$, the number of viruses inhaled during an outing $I_V$ (virions), which is considered to have a direct correlation with infection, could be used as a parameter for evaluation. If the estimated viral concentration $d$ (virions/m$^3$) in the activity range is available, the following definition can be adopted.

$$I_V = \text{number of viruses inhaled during outings (virions)}$$

$$= \int [d (1 - R_{S,in}) Q_{\text{breath,in}}] \, dt$$

where, $d$ (virions/m$^3$) = estimated viral concentration at the location, $R_{S,in}$ = aerosol shielding ratio for the air supply of PAPR, and $Q_{\text{breath,in}}$ (m$^3$/s) = estimated amount of inhaled air of the wearer at the time (exhaled air is not counted). As a request from the government to each citizen, it is assumed that keeping the above $W_R$, $T_{WT}$, or $I_V$ at a certain level or better will be requested.

**Ethical Considerations**

This study is based on known facts and the author's own thinking, and no new experiments were performed. Therefore, the author has not applied to Gunma University, to which the author belongs, for ethics approval. Consent for publication has been granted from the identifiable individual (author YF) in Figure 1 in this paper.

**Results**

**What PAPR Can Do as an Alternative to Lockdown**

PAPR was shown to be a possible alternative to lockdown through the reduction of airborne and droplet transmissions and through the temporary reduction of $\beta$. The existing medical PAPRs appear to have sufficiently high virus shielding performance and appear to have already reached a level that should be experimentally tested as an alternative to lockdown. The current medical PAPR shown in Figure 1 is expensive and does not have measurement control functions. However, the prototypes shown in Figure 1 indicate that cost reduction and high functionality are possible. In addition, a variety of PAPRs are commercially available for nonmedical use, some of which are inexpensive. If an inexpensive PAPR is supplied to everyone, using PAPRs during a pandemic instead of issuing a countrywide lockdown will become a reality.
How to Operate a Social System Utilizing PAPR

A realistic process is shown below for quantitatively evaluating the effect of the aerosol shielding performance of PAPRs (for air intake side and exhaust side), PAPR wearing rate and wearing condition for reducing the β, and the effective reproduction number $R_t$ for realizing this proposal.

1. Select and prepare special experimental zones for social experiments in the next pandemic.

2. In the experimental zone, when lockdown is applied to the surrounding area, PAPR can be substituted for the various activity restrictions during lockdown.

3. Compare the spread of infection between the special experimental zone and other areas, and change the aerosol shielding performance of PAPR (air supply side and exhaust side), PAPR wearing rate and condition, and other operational conditions within the special experimental zone. The obtained results can be used to quantitatively evaluate the effects of the aerosol shielding performance of PAPR (air supply side and exhaust side), PAPR wearing rate and wearing condition for reducing the β, and the effective reproduction number $R_t$.

4. When the effectiveness of PAPR as an alternative to lockdown is confirmed and the problems are sufficiently resolved, PAPR as an alternative to lockdown can be applied to other regions. Examples of operations in special experimental zones include the following: (1) people can go out freely if they wear PAPRs, even in circumstances where going out is restricted in other surrounding areas and (2) factories can be operated freely if its employees wear PAPRs, even in circumstances where factories are prohibited to operate in other surrounding areas. Figure 4 shows the proposed social system where PAPRs are used as an alternative to lockdown.

In the proposal in this research, PAPR will be utilized under the leadership of the government as described below:

1. The government distributes PAPRs (helmet/hood type) to all citizens as emergency equipment.

2. If the estimated effective reproduction number $R_t$ is high and there is concern about an outbreak of infection, the government will determine the type and intensity of the lockdown and decide how to replace each constraint during lockdown with PAPR. Examples include prohibition on going outside (going outside is possible if wearing a PAPR), prohibition of factory operation (operation is possible if all employees wear PAPRs), prohibition of restaurant operation (operation is possible if all employees wear PAPRs and all customers wear PAPRs suitable for eating and drinking), and overseas entry prohibition (entry is allowed if visitors agree to wear a negative pressure PAPR for a specified period of time and accept government remote monitoring of wearing conditions). This will make it possible to open the door to foreigners and returnees who wish to enter the country, although they would be subject to the same level of inconvenience as ordinary citizens suspected of being infected.

In the initial implementation of the proposed social system, as described above, special experimental zones will be established in various regions, various trials will be conducted based on various assumptions, and data will be collected. Based on the data obtained, qualitative and quantitative evaluations of the benefits (reduction of infection probability) and burdens borne by individuals and the benefits (reduction of infection spread) and burdens for the society as a whole will be attempted. The proposed social system should be compared and verified with the lockdown in each of the different situations, and the best way to be found as an alternative to the various restrictions.
imposed by the lockdown should be identified. Ultimately, a PAPR-utilizing social system will be constructed that effectively functions as an alternative to lockdown.

**Directions for Improvement of PAPRs as an Alternative to Lockdown**

The following 7 points can be considered as directions for improvements of PAPRs as a lockdown alternative.

**Flow Path Optimization**

A hood shape and part configuration should be developed that provides a smooth flow of the exhaled air out of the hood. The concentration of carbon dioxide is approximately 500 ppm (0.05%) in ambient air and approximately 50,000 ppm (5%) in exhaled air [30]. The oxygen concentration in the exhaled air is expected to decrease from the oxygen concentration in the ambient air (approximately 21%) by an amount equal to the increase in the carbon dioxide concentration in the exhaled air (approximately 5%). In the commercially available PAPR and the developed PAPRs shown in Figure 1, a large flow rate (approximately 200-400 L/min) is delivered compared to the resting respiratory flow rate (approximately 6-10 L/min) [31] in order to suppress the carbon dioxide concentration in the hood [25,26,29]. Efficient expiration of exhaled air to the outside allows for a significant reduction in the air supply flow rate, resulting in a significant reduction in the size and weight of pumps, batteries, and filters, as well as design diversification.

By minimizing the volume inside the mask, it is also possible to minimize the retention of the exhaled air from the nose and mouth inside the mask. As an extreme example, consider a configuration in which the nose is used for inhalation, the mouth is used for exhalation, and the air supply to the nose and the exhaust from the mouth are mechanically separated. As a result, the flow rate of the air filtered through the nonwoven filter and delivered to the nose becomes the same as the flow rate from the nose, and this dramatic reduction in flow rate results in a drastic reduction in the pump and battery capacity.

**Precise Pressure Control by Fluid Modeling**

Fluid modeling of PAPRs should be considered. For the PAPR prototypes (simple PAPR and controller PAPR) shown in Figure 1, the air supply through a nonwoven fabric filter is realized by a pump, and the exhaust through a nonwoven filter is created through the positive internal pressure. A simple modeling for these PAPRs is as follows.

**Air Supply Flow Rate**

The flow rate \( Q_{in} (\Delta P, V) \) through the filter is determined by the pressure difference \( \Delta P \) before and after the filter. The flow rate through the pump is determined by the pressure difference \( \Delta P \) before and after the pump and the applied voltage \( V \) of the pump. When the differential pressure \( \Delta P (=\Delta P_t + \Delta P_p) \) inside and outside the PAPR and the pump applied voltage \( V \) are determined, the air supply flow rate \( Q_{in} \) is determined.

**Exhaust Flow Rate**

The flow rate \( Q_{out} (\Delta P) \) through the filter is determined by the pressure difference \( \Delta P_t \) before and after the filter.

**Respiratory Flow**

The flow difference \( Q_{diff} \) between the air supply flow rate \( Q_{in} \) and the exhaust flow rate \( Q_{out} \) can be expressed as follows.

\[
Q_{diff} = Q_{in} (\Delta P, V) - Q_{out} = Q_{breath} + Q_{leak} + Q_{volume}
\]

Here, \( Q_{breath} \)=respiratory flow rate of the wearer of PAPR, positive with inspiration; \( Q_{leak} \)=leak flow rate, positive for leakage from the inside to the outside; and \( Q_{volume} \)=volume change inside PAPR, positive with volume increase.

Since the time averages of respiratory flow \( Q_{breath} \) and volume change \( Q_{volume} \) are zero, the time average of \( Q_{diff} \) is the time average of \( Q_{leak} \). In addition, \( Q_{leak} \) is expressed as a function \( Q_{leak} (\Delta P) \) of the differential pressure \( \Delta P \), assuming that the shape of the gap between the face and the mask is constant. Furthermore, the volume change \( Q_{volume} \) is considered to be expressed as a function \( Q_{volume} (\Delta P) \) of the differential pressure \( \Delta P \).

\[
Q_{breath} = Q_{diff} - Q_{leak} (\Delta P) - Q_{volume} (\Delta P)
\]

In this case, \( Q_{breath}<0 \) is judged as expiration, and \( Q_{breath}>0 \) is judged as inspiration. In this way, the exhalation and inhalation movements of the wearer can be detected in real time. For example, based on this detection result, the following control can be considered.

1. If an exhalation movement is detected, the minimum positive pressure setting (eg, 10 Pa) is set to minimize the resistance to exhalation movement while preventing leakage from the gap.

2. If an inhalation movement is detected, a strong positive pressure setting (eg, 100 Pa) is used to positively assist the inhalation movement.

In addition to the forced air supply by the pump and filter, the introduction of forced exhaust by the pump and filter enables the following differential pressure control.

1. When an exhalation movement is detected, a strong negative pressure setting (eg, –100 Pa) is used to actively assist the exhalation movement.

2. When an inhalation movement is detected, a strong positive pressure setting (eg, 100 Pa) is used to actively assist the inhalation movement.

In light of this, a PAPR facilitates the easy movement of the wearer’s exhalation and inhalation. In this case, the direction of leak flow at the possible gap is opposite to normal—from outside to inside during exhalation and from inside to outside during inhalation. It will be possible to detect coughing from the measurement results of the differential pressure \( \Delta P \). It will also be possible to estimate the possibility of infection of the wearer together with other measurement results such as body temperature. The ability to efficiently identify infected persons will enable efficient isolation and treatment of infected persons and will have a significant effect in reducing the spread of infection throughout the society.
Improved Design

All the 3 types of PAPRs shown in Figure 1 have bulky and exaggerated designs. As a lockdown alternative, the design may not be very important; however, it is better to have an excellent design. If the above flow path optimization achieves a dramatic reduction in the air supply flow rate, then a dramatic reduction in pump and battery size and various designs will become possible. Once PAPRs are widely accepted as a lockdown alternative, many people will be dissatisfied with the bare-bones PAPRs provided by the government; it is conceivable that companies of various genres will develop models with different characteristics.

Maintenance Method

It is necessary for every citizen to be able to easily perform maintenance such as cleaning and disinfecting the PAPR and replacing the nonwoven filter unit at home. However, when PAPRs are used as an alternative to lockdown, it is expected that the virus concentration in the external environment will be extremely low compared to the environment assumed in medical PAPRs due to the following reasons.

1. Infected persons would wear PAPRs.
2. PAPR has the ability to not only stop the entry of droplets and aerosols containing viruses but also prevent their release to the outside.
3. PAPR purifies indoor air in the same way as an air purifier.

Therefore, in terms of maintenance standards, it may be possible to set relatively lenient standards for nonmedical use PAPRs compared to those set for medical PAPRs, which are assumed to be used in environments with high virus concentrations such as hospital wards where infected people are congregated.

Variations Suitable for Different Places of Use and Activity Contents

The following variations should be developed, which are fine-tuned to suit various places of use and activities, as well as to suit societies and populations at different stages of social, economic, and cultural development: (1) a model that pursues comfort for everyday use; (2) models suitable for specific activities such as sports, eating, and drinking, for example, a model for eating and drinking with a face shield opening and closing mechanism and an air shower function or a model for jogging with a structure that mechanically separates the nose (exhalation) and mouth (inhalation); (3) a booth type model that wraps around a desk and a chair in an office, vehicle, restaurant, etc; (4) a model compatible with the standard unit of the ceiling-mounted air conditioner; and (5) a very inexpensive model suitable for low-income countries and regions.

PAPR With Information Terminal Function

The PAPR prototype (controller type) shown in Figure 1 is an all-in-one type PAPR with a computer and a power supply on the wearer’s head. Therefore, it is easy to make the PAPR an advanced information terminal by means of installing a computer equivalent to that of a high-end smartphone and adding various devices as follows: (1) equipped with smartphone function and virtual reality screen, (2) equipped with a noncontact input system using eye gaze and brain waves, and (3) equipped with a physical condition measurement and management system using body temperature sensors, cough sensors (pressure sensors), electroencephalogram sensors, etc. If the PAPR is comfortable to breathe, comfortable to wear, and has advanced information terminal functions, it is expected that some people will not be able to part with it. In particular, if a physical condition measurement and management system is installed to accurately estimate the presence or absence of infection, it will be easier to isolate, examine, and treat those who are deemed to have a high probability of being infected. In many cases, the various behavioral and activity restrictions during lockdown are uniformly applied to all persons under conditions where it is not known who is infected. If it becomes possible to know with a high degree of certainty who is infected, the use of PAPR as an alternative to lockdown can be changed to a more targeted approach.

Evaluation Indicators and Evaluation Methods

As an evaluation indicator of PAPR as a lockdown alternative, it is desirable to be able to quantitatively evaluate the effect of reducing the aforementioned \( \beta \) by means of wearing a PAPR. However, in order to estimate the \( \beta \) reduction rate with high accuracy, it is necessary to conduct elemental experiments and social experiments under various conditions.

The most important evaluation indicators of the basic performance of PAPR that should be obtained from elementary experiments are as follows: (1) reduction rate of virus-containing aerosols and droplets inhaled by potentially infectious persons wearing PAPR and (2) reduction rate of virus-containing aerosols and droplets exhaled by infected persons wearing PAPR. Current standards (eg, APF) usually refer only to the reduction rate of virus-containing aerosols and droplets inhaled by potentially infectious persons wearing PAPR. However, both (1) and (2) are considered to be equally important when requiring the uniform wearing of PAPRs by the general public in cases where presence or absence of infection is unclear for the purpose of reducing the effective reproduction number \( R_e \).

In the 3 PAPR models shown in Figure 1, positive pressure was used to prevent outside air from entering directly through gaps, with an emphasis on protecting the inside (wearer) from the outside. If the above (1) and (2) are equally important, then it is equally important to protect the wearer from the outside environment and to protect the people from the wearer, thereby indicating the it is not necessary to make the internal pressure positive.

Impact of the PAPR Internal Environment on the Mind and Body

If PAPR is considered as an alternative to lockdown measures, the impact of the PAPR internal environment on the mind and body of the wearer will become important. It is necessary to comprehensively investigate the relationship between the following 2 types of parameters from the viewpoint of the influence of the PAPR internal environment on the mind and body of the wearer.
Physical Parameters Related to the PAPR Internal Environment

The physical parameters to be considered are concentrations of particulate pollutants (droplets, aerosols, pollen, particulate matter 2.5, mite corpses, dust, etc), gaseous pollutants, gas composition (carbon dioxide concentration, oxygen concentration, etc), differential pressure, temperature, humidity, acoustic characteristics (sound transfer characteristics, noise, etc), vibration, and airflow.

Biological and Psychological Parameters of PAPR Wearsers

The biological and psychological parameters are respiratory status, electroencephalogram, body temperature, pulse rate, comfort, safety (physical danger, probability of infection), and degree of relaxation.

Balance Between Infection Control Efficiency and Personal Freedom Through the Use of IoT

As shown in Figure 5, the operation of the PAPR wearing rate network management system led by the government will be performed as follows.

Figure 5. The social system with the powered air purifying respirator wearing rate network management system.

<table>
<thead>
<tr>
<th>Normal State (No Concern of Infection Explosion)</th>
<th>Concern of Infection Explosion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present Social System</td>
<td>Small</td>
</tr>
<tr>
<td>New lifestyle + vaccination</td>
<td>Large</td>
</tr>
<tr>
<td>Status of Lockdown</td>
<td>Weak</td>
</tr>
<tr>
<td>Required Wearing Rate ($W_{R,required}$) for everyone</td>
<td>0%</td>
</tr>
<tr>
<td>Proposed Social System</td>
<td>100%</td>
</tr>
</tbody>
</table>

1. The government will distribute PAPRs (with smartphone connectivity) with sufficient aerosol shielding performance to all citizens as emergency equipment. Each citizen installs an app with a wearing rate proof function on his/her own smartphone.

2. If the estimated effective reproduction number $R_e$ is high and there is concern about an outbreak of infection, the government will (1) set the target effective reproduction number $R_{e,target}$, (2) solve the formula based on appropriate assumptions to calculate the required wearing rate $W_{R,required}$ required to achieve $R_{e,target}$, (3) show $W_{R,required}$ and require all citizens to comply with it—each citizen can spend their time without PAPR at any place (party venue, restaurant, pub, etc) and any time by showing proof of wearing rate $W_R$ within the scope of fulfilling their obligations, and (4) pay close attention to changes in the effective reproduction number $R_e$ and raise $W_{R,required}$ if the goal of controlling the spread of infection is in jeopardy. Conversely, if it exceeds the target, lower $W_{R,required}$ and increase the degree of freedom in citizen life.

If the PAPR lockdown alternatives are strong enough, the government can quickly contain the spread of infection by setting $W_{R,required}$ to 1.0 (100%) even when the government makes a big mistake in estimating $W_{R,required}$ and falls into the worst situation. In that case, the government can conduct various trials and countermeasures on various assumptions and hypotheses with a leeway. The government is freed from constraints that limit them to overly conservative measures. In addition, throughout the entire process, the government will be able to improve the accuracy of the above formula based on appropriate assumptions by using the big data collected on the relationships between “changes in aerosol shielding performance of PAPR (air intake side, exhaust side), PAPR wearing rate $W_R$, and wearing condition, etc” and “changes in infection spread status and the effective reproduction number $R_e$.”

Discussion

Principal Findings

In this study, the feasibility of the following 2 ideas was examined. First, the construction of a social system using PAPR...
with similar infection control ability as lockdown measures and with less economic and social damage as an alternative to lockdown is possible. Second, balancing the efficiency of the government’s infection control and each citizen’s personal freedom is possible by means of an IoT system.

**Extended Functionality and Privacy Protection in the PAPR Wearing Rate Network Management System**

By utilizing PAPR with several sensors (thermometer, cough sensor, etc), the government can make this system much more powerful than conventional apps for measuring contact with infected persons. For example, the system may be able to improve the accuracy of infection detection based on big data concerning changes in body temperature, cough (condition and frequency), and the presence or absence of severe disease after infection until the onset of illness. In the case of PAPR equipped with both air supply and exhaust pumps, the wearer can switch the internal pressure between positive pressure when not infected and negative pressure when infected, thus prioritizing the prevention of the spread of infection in the society as a whole. From the viewpoint of privacy protection, social discussion is necessary for the following matters: (1) how much of the information from PAPRs should be passed on to the government server? and (2) how should the government’s use of personal information be curbed? Especially for (2), it is considered necessary to develop and construct a technical and social mechanism to realize a brake. In order to prevent misuse of personal information, it is conceivable to apply the proposal for a street camera system’s perfect recording of usage history by a reliable third party as the first step [30]. Methods to substitute each restriction in lockdown with PAPR utilization need to be considered in various social systems. Those who wish to use PAPR as a substitute for the constraints imposed by lockdown need to prepare to obtain the PAPR before the pandemic. Therefore, it is also important for the society as a whole to ensure and disseminate information on how to obtain PAPRs.

Different countries have different governance systems. In some countries, it might not be easy to make the public understand that PAPR can be used as an alternative to lockdown measures—they may make it an option and not a mandate. This paper discusses how PAPR can substitute the primary constraints imposed by lockdown. Even in cases of other alternatives such as combination of lockdown and free mobility of low-risk populations during COVID-19 [32], PAPRs may be used for controlling the infection rates. The proposed PAPR wearing rate network management system utilizes IoT technology, which is currently being widely pursued by various societies and companies. In order to build a social system that makes the government’s control of the spread of infection more efficient and that respects the freedom of individuals to the maximum extent, social experiments should be first conducted under various conditions to identify the challenges and improve the effectiveness of PAPRs.

**Society of People Breathing Purified Air**

Further, although this is a discussion that is far from the main point of this paper, if truly high-performance, comfortable, and low-cost PAPRs are successfully developed through this research and subsequent research and developments, it is possible that many people will desire PAPR-purified air instead of the air around them. This is similar to the situation of drinking water, that is, just as how populations consume purified water through water-treatment and water-purification technology rather than water from ponds and rivers, people would prefer purified air to breathe. It can be expected that many citizens will wear PAPRs when they go out, regardless of whether the government asks them to do so. As more people breathe purified air, there may be concerns about the public’s immune system being weakened against airborne diseases and pollen allergies. However, there is no dispute that water-borne infectious diseases have become controllable because many people drink only purified water, and it would not be advisable to drink water without purification, as was the case in primitive times. A society in which the majority of the population breathes purified air will be resilient to all airborne diseases. The construction of such a society has the potential to be an opportunity for a historic change in the human race, which has been plagued by airborne diseases.

**Conclusions**

This study examines the feasibility of 2 ideas. First, this study shows that it is possible to construct a social system using PAPR with similar infection control effects as lockdown measures and with less economic and social damage as a means of temporarily reducing the effective reproduction number $R_e$. Second, the PAPR wearing rate network management system balances the achievement of the efficiency of the government’s infection control and each citizen’s personal right to choose the time and opportunity not to wear PAPR during a pandemic.

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**Data Availability**

This study was conducted using deductive thinking based on publicly available information. No confidential data were used in this study.


19. Assigned protection factors (APF) for 3M hoods and helmets. 3M 2021. URL: https://tinyurl.com/4m33pn4h [accessed 2024-04-28]


Abbreviations

APF: assigned protection factor
IoT: Internet of Things
PAPR: powered air purifying respirator

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Stroke Survivors’ Interaction With Hand Rehabilitation Devices: Observational Study

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Abstract

Background: The hand is crucial for carrying out activities of daily living as well as social interaction. Functional use of the upper limb is affected in up to 55% to 75% of stroke survivors 3 to 6 months after stroke. Rehabilitation can help restore function, and several rehabilitation devices have been designed to improve hand function. However, access to these devices is compromised in people with more severe loss of function.

Objective: In this study, we aimed to observe stroke survivors with poor hand function interacting with a range of commonly used hand rehabilitation devices.

Methods: Participants were engaged in an 8-week rehabilitation intervention at a technology-enriched rehabilitation gym. The participants spent 50-60 minutes of the 2-hour session in the upper limb section at least twice a week. Each participant communicated their rehabilitation goals, and an Action Research Arm Test (ARAT) was used to measure and categorize hand function as poor (scores of 0-9), moderate (scores of 10-56), or good (score of 57). Participants were observed during their interactions with 3 hand-based rehabilitation devices that focused on hand rehabilitation: the GripAble, NeuroBall, and Semi-Circular Peg Board. Observations of device interactions were recorded for each session.

Results: A total of 29 participants were included in this study, of whom 10 (34%) had poor hand function, 17 (59%) had moderate hand function, and 2 (7%) had good hand function. There were no differences in the age and years after stroke among participants with poor hand function and those with moderate (P=.06 and P=.09, respectively) and good (P=.37 and P=.99, respectively) hand function. Regarding the ability of the 10 participants with poor hand function to interact with the 3 hand-based rehabilitation devices, 2 (20%) participants with an ARAT score greater than 0 were able to interact with the devices, whereas the other 8 (80%) who had an ARAT score of 0 could not. Their inability to interact with these devices was clinically examined, and the reason was determined to be a result of either the presence of (1) muscle tone or stiffness or (2) muscle weakness.

Conclusions: Not all stroke survivors with impairments in their hands can make use of currently available rehabilitation technologies. Those with an ARAT score of 0 cannot actively interact with hand rehabilitation devices, as they cannot carry out the hand movement necessary for such interaction. The design of devices for hand rehabilitation should consider the accessibility needs of those with poor hand function.

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KEYWORDS

stroke; rehabilitation; hand rehabilitation devices; accessibility; stroke survivors; rehabilitation technologies
Introduction

Stroke is a major cause of disability in the world [1]. Globally, about 17 million people have a stroke each year [2]. In the United Kingdom, the prevalence of stroke is projected to rise from 950,200 to 2,119,400 cases between 2015 and 2035 [3]. This projected rise in the prevalence of stroke has been associated with improvements in medical advances that have led to a decline in the number of deaths due to acute stroke, among other reasons [4]. Nevertheless, stroke survivors are faced with considerable long-term periods of enduring physical impairments, the likelihood of reoccurrence of strokes, transient ischemic attacks, or even death within 1 year of having a stroke [5]. Motor impairment (muscle weakness and the loss of movement control) is the most common consequence of stroke, impacting several aspects of life and reducing the ability of stroke survivors to lead an independent life [6]. About 55% to 75% of those who survive a stroke experience motor impairment in the upper limb 3 to 6 months after stroke [7].

The hand is crucial for carrying out activities of daily living such as eating, dressing, bathing, and communicating [8]. Besides, the hand is a defining feature of human beings and is vital for human daily interaction [9]. Due to this importance, impairments such as spasticity and weakness, which are common sequelae of stroke [10] and manifest in a fixed flexed position of the wrist and fingers, affect the function of the hand and impact the quality of life [10].

Rehabilitation can have a positive impact on the recovery of functions in persons with stroke [11] as well as in enhancing their quality of life [12], and movement restoration is a key goal in the rehabilitation of persons with neurological disorders [13]. The relearning of movement ability during rehabilitation is based on factors such as the repetitiveness, intensity, and regularity of task-specific movements [14]. It has been suggested that the rehabilitation of hand mobility and strength be prioritized once the general physical situation of stroke survivors has been stabilized owing to the importance of the hand [15].

Several new rehabilitation technologies that target the upper limb to improve motor functions are currently in use; these include the use of robotic-assisted technologies, virtual reality, and telerehabilitation [16]. Some others that are used in this study are gaming devices such as the GripAble (Gripable), NeuroBall (Neurofenix), and Semi-Circular Peg Board (Rolyan). The NeuroBall is an interactive device that connects wirelessly with a tablet app to carry out activities that can also be objectively measured [17]. The GripAble is a similar lightweight electronic handgrip [18] that also interacts wirelessly with a computer tablet, enabling users to interact with therapy games tailored to improve the upper limb and hand function in a way that can be objectively assessed [18,19]. The Rolyan Semi-Circular Peg Board consists of 3 colored pegs (red, white, and blue) of different diameters that the users are expected to pick up and place in their different peg holes (based on their diameter; see Figure 1 below). The ability of stroke survivors with poor hand function to access these devices is a major concern, as according to a report [20], only hemiplegic stroke survivors who are mildly disabled are likely to access hand or arm training apps that are available on mobile devices.

This study aims to observe stroke survivors’ interaction with hand rehabilitation devices and to understand how the different categories of hand function (Action Research Arm Test [ARAT] scores) influence the stroke survivors’ rehabilitation goals.
Methods

Participants

Participants were recruited from cohorts of stroke survivors attending a rehabilitation intervention at a cocreation center for accessible rehabilitation technology [21] between September 2021 and April 2023. The inclusion criteria for this study have been described in detail previously [21]; briefly, participants had to have had a stroke within the last 12 months that resulted in mobility problems, be aged over 18 years, be well enough to engage in light to moderate exercise, and be able to attend the rehabilitation program at least twice a week. A range of outcome measures were taken before and after the program, including the ARAT. An overview of the full rehabilitation program is available in our previously published report [21].

Out of a total of 36 participants who agreed to take part in the intervention, 7 (19%) were excluded from this study. Of the 7 excluded persons, 5 (71%) withdrew from the intervention (2/5, 40% withdrew before the commencement and 3/5, 60% withdrew due to ill health or unwillingness to continue), and the other 2 (29%) of the 7 were excluded as a result of incomplete data.

The Upper Limb Rehabilitation Intervention

The upper limb intervention involved activities designed to improve the upper limb functions of participants, delivered completely through the use of technology and therapy devices that either stimulated or promoted repetitive and intensive movement training. The upper limb and hand rehabilitation technologies available to the participants in this study are shown in Table 1. The participants spent at least 50-60 minutes of each of the 2-hour sessions engaging with these devices.
Table 1. Upper limb rehabilitation technologies used.

<table>
<thead>
<tr>
<th>Technology or device</th>
<th>Manufacturer</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>GripAble</td>
<td>Gripable</td>
<td>It connects wirelessly with an app on a computer tablet [19] to interact with specifically designed therapy games [22], to train 4 different types of upper limb movements, such as grip and release, pronation and supination, wrist flexion and extension, and radius and ulnar deviations.</td>
</tr>
<tr>
<td>NeuroBall</td>
<td>Neurofenix</td>
<td>It connects wirelessly with a tablet app and interacts with therapy games specifically designed to exercise the upper limb of stroke survivors [17]. It trains upper limb movements such as finger grip; hand grip; right, left, upward, and downward tilt; and elbow and shoulder movements.</td>
</tr>
<tr>
<td>Mirror box</td>
<td>Saebo</td>
<td>It is a form of mental practice that excites the primary motor cortex, thereby evoking the movement of the affected limb, as the participants move the unaffected side while looking into the mirror [23].</td>
</tr>
<tr>
<td>Sensory TENS(^a)</td>
<td>Med-Fit</td>
<td>It is a noninvasive nerve stimulator used to relieve pain [24], stimulate the muscles, and relieve muscle stiffness [25].</td>
</tr>
<tr>
<td>Semi-Circular Peg Board</td>
<td>Rolyan</td>
<td>It is a therapy tool designed to improve upper limb strength, movement coordination, endurance, and range of motion. It aims to improve hand dexterity.</td>
</tr>
<tr>
<td>Armeo Spring</td>
<td>Hocoma</td>
<td>It provides arm weight support while encouraging users to carry out self-initiated arm movements in the shoulder, elbow, and wrist joints and trains different upper limb movements [26].</td>
</tr>
<tr>
<td>Vibrating or hot compress massage ball</td>
<td>Dongguan Kooeej</td>
<td>It stimulates the hand using the vibrations delivered at different intensities.</td>
</tr>
<tr>
<td>VR(^b) headset</td>
<td>Oculus Quest with In-cisiv software</td>
<td>It immerses the user into a virtual environment, thereby encouraging them to use their affected limb to interact with functional tasks [27, 28].</td>
</tr>
</tbody>
</table>

\(^a\)TENS: transcutaneous electrical nerve stimulation.
\(^b\)VR: virtual reality.

Overview of the Upper Limb Rehabilitation Program

Figure 2 is a representation of the upper limb rehabilitation program used in the rehabilitation gym. The activities were divided into 2 categories. The first part aimed at priming the brain to prepare it for plastic response [29]. Priming focused on sensory stimulation including mirror therapy and electrical, thermal, and vibrational stimulation. These priming activities comprised the first 15-20 minutes of each rehabilitation session. This second part, that is, the “active training,” aimed to engage the participants in high-intensity motor tasks such as object grip and release, object manipulation, and reach to grasp, designed to improve range of motion, strength, and control. The participants were not limited in terms of the number of devices they could use.

Figure 2. Upper limb rehabilitation program model for stroke survivors. TENS: transcutaneous electrical nerve stimulation.

Categorizing Participants Into Different Hand Function Groups

Participants were given a 1-day initial appointment with a therapist at the rehabilitation gym before the commencement of the 8-week rehabilitation intervention. During this appointment, demographic data including stroke history were collected, along with a range of baseline assessments for mobility, communication, and cognition, including the ARAT [30]. The ARAT was used to categorize the participants into 3 different hand function groups: poor (scores of 0-9), moderate (scores of 10-56), and good (score of 57) [30].
Understanding the Rehabilitation Goals of Those With Different Categories of Hand Function

During the preintervention visit, participants were allowed to communicate their rehabilitation goals and interact with the upper limb devices to understand how they are set up and operated. The rehabilitation goals of the participants were summarized based on their different hand functions to help understand the needs of stroke survivors who fall under each of the different hand functions, particularly the hand rehabilitation goals of those with poor hand function.

Observing the Interaction of Those With Poor Hand Function and the Hand Rehabilitation Devices

Following the goal setting and initial interaction with the devices, a rehabilitation program was drawn up. The rehabilitation program was individually tailored by a physiotherapist using the rehabilitation goals of the participants. The program however only acted as a guide, as participants had the freedom to interact with any of the devices. The ability of the participants to use each rehabilitation device was observed and recorded. At the end of the intervention, all the observations from participants with poor hand function were gathered and studied to see how they interacted with the hand-based rehabilitation devices. Three of the upper limb devices—the GripAble, NeuroBall, and Semi-Circular Peg Board (see Figure 1)—were selected for observation in this study. The reason for selecting these devices is because these 3 devices were the only devices listed under the “active training” category (see Figure 2) at the time of the study that were used to primarily train motor activities in the hand (involving the wrist and fingers) in addition to training other parts of the upper limb.

Data Organization and Analysis

The simple percentage method was used to estimate the percentage of stroke survivors who fall into each category of hand function. A 1-way ANOVA was carried out using Minitab statistical software (Minitab LLC), with the Dunnett multiple comparison method used to compare the ages of the group with poor hand function to those with moderate and good hand function.

Ethical Considerations

This study was approved by the University of Strathclyde ethics committee (approval UEC 20/08). The participants provided written informed consent before the study, and their participation was voluntary (no compensation was provided). All identifiable data were pseudonymized and replaced with a code.

Results

Categorizing Participants Into Different Hand Function Groups

Observations from 29 participants were included in this study. Their average age was 59.10 (SD 13.62) years with an average of 3.140 (SD 2.31) years after stroke. Of the 29 participants, 17 (59%) were hemiplegic on the left side of their body, whereas the remaining 12 (41%) were hemiplegic on the right side of their body (Table 2).

Of the 29 participants, 10 (34%) scored between 0 and 9 on the ARAT and were grouped as having poor hand function, 17 (59%) scored between 10 and 56 on the ARAT and were grouped as having moderate hand function, and 2 (7%) scored 57 on the ARAT and were grouped as having a good hand function. There was no statistical difference in age between the poor hand function group and both the moderate hand function \((P=0.06)\) and the good hand function \((P=0.37)\) groups. Similarly, there was equally no difference in the years after a stroke between the poor hand function group and both the moderate hand function \((P=0.09)\), and the good hand function \((P=0.99)\) groups. There was also no observed difference in the hemiplegic side of those with poor hand function (left: 5/10, 50%; right: 5/10, 50%).

Table 2. Characteristics of participants and the 3 subgroups.

<table>
<thead>
<tr>
<th>Group</th>
<th>Participants (n=29), n</th>
<th>Hand function</th>
<th>Age (years), mean (SD)</th>
<th>Years after stroke, mean (SD)</th>
<th>Hemiplegic side, n (%)</th>
<th>ARAT(^a) score, mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All</td>
<td>29 (100)</td>
<td></td>
<td>59.10 (13.62)</td>
<td>3.14 (2.31)</td>
<td>17 (59)(^c)</td>
<td>26.63 (21.51)</td>
</tr>
<tr>
<td>1</td>
<td>10 (34)</td>
<td>Poor</td>
<td>64.70 (8.83)</td>
<td>2.10 (1.45)</td>
<td>5 (50)(^d)</td>
<td>2.00 (3.74)</td>
</tr>
<tr>
<td>2</td>
<td>17 (59)</td>
<td>Moderate</td>
<td>53.76 (13.89)</td>
<td>3.88 (2.57)</td>
<td>11 (65)(^e)</td>
<td>34.65 (16.09)</td>
</tr>
<tr>
<td>3</td>
<td>2 (7)</td>
<td>Good</td>
<td>76.50 (0.707)</td>
<td>2.00 (1.42)</td>
<td>1 (50)(^f)</td>
<td>57.00 (0.00)</td>
</tr>
</tbody>
</table>

\(^a\)ARAT: Action Research Arm Test.
\(^b\)Not applicable.
\(^c\)n=29.
\(^d\)n=10.
\(^e\)n=17.
\(^f\)n=2.
Understanding the Rehabilitation Goals of Those With Different Categories of Hand Function

Table 3 shows a summary of the rehabilitation goals of stroke survivors based on their different hand functions. Participants with poor hand function stated goals that were more toward gaining movements in different parts of their upper limb, as well as improving the ability to carry out active movements that will enable them to grasp and release objects. However, stroke survivors with moderate and good hand function had goals that were focused on how to improve grip strength, fine motor movements, release time, as well as purposeful movement of the upper limb (see Table 3). Those with poor hand function who recorded a score greater than 0 on the ARAT equally communicated the need to improve grip strength.

Table 3. Upper limb and hand rehabilitation goals of participants separated into the 3 functional categories.

<table>
<thead>
<tr>
<th>Group</th>
<th>Hand function</th>
<th>Rehabilitation goals as stated by the participants</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Poor</td>
<td>• Gain the ability to hold objects (e.g., paper)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Gain some shoulder movement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Gain arm movement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Recovery of any movement, primarily in the shoulder</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve the grasp and release of objects</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve active movements</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Grip strength&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>2</td>
<td>Moderate</td>
<td>• Improve dexterity</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve grip</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve the range of upper limb movement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve upper limb strength</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve supination or pronation range</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve the grasp and release of objects</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve release time</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Gain the ability for small object manipulation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Gain the ability to move objects</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Gain the ability for purposeful movement of the upper limb</td>
</tr>
<tr>
<td>3</td>
<td>Good</td>
<td>• Increase grip</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Improve wrist extension</td>
</tr>
</tbody>
</table>

<sup>a</sup>For those who recorded a score >0 on the Action Research Arm Test (ARAT).

Interaction With Hand Rehabilitation Technologies by the Poor Hand Function Group

Table 4 shows that 8 (80%) of the 10 participants with poor hand function could not interact with any of the 3 aforementioned devices to carry out active training. This value represents 28% (8/29) of the total population in this study. Only 2 (20%) of the 10 participants with poor hand function were able to engage with these devices; the ARAT score shows that these 2 participants had ARAT scores of 7 and 9, compared to the score of 0 that was recorded by the other 8 who were not able to engage with these devices.
Table 4. Interaction of stroke survivors who had poor hand function with the hand rehabilitation devices.

<table>
<thead>
<tr>
<th>Participant ID</th>
<th>ARAT score</th>
<th>Upper limb rehabilitation goal</th>
<th>Use of devices for active hand training</th>
<th>Comments on the participants’ ability to use the devices</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>GripAble</td>
<td>NeuroBall</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>General upper limb function</td>
<td>✓</td>
<td>X</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>Improve active movements</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>3</td>
<td>0</td>
<td>Hold objects (eg, paper), gain some shoulder movement</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>Improve the grasp and release of object</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>5</td>
<td>7</td>
<td>Grip strength, range of shoulder or elbow active movement</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>6</td>
<td>0</td>
<td>Gain arm movement</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>7</td>
<td>0</td>
<td>Would like to get some movement</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>8</td>
<td>0</td>
<td>Recovery of any movement, primarily in the shoulder</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>9</td>
<td>0</td>
<td>—d</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>10</td>
<td>9</td>
<td>Grip strength</td>
<td>✓</td>
<td>✓</td>
</tr>
</tbody>
</table>

ARAT: Action Research Arm Test.

X: unable.

✓: able.

d: Not applicable.

Discussions

Principal Findings

This study was carried out to observe how stroke survivors with poor hand function interacted with hand rehabilitation devices such as the GripAble, NeuroBall, and Semi-Circular Peg Board. The findings show that stroke survivors whose poor hand function leads to an ARAT score of 0 cannot actively interact with hand rehabilitation devices.

Comparison to Prior Work

About two-thirds (55%-75%) of persons who had a stroke sustain upper limb impairments [7]. The extent of the impairments varies from person to person (see Table 2). In some, it results in poor hand function, whereas others present moderate or good hand function. The level of hand function present after stroke subsequently influences the upper limb rehabilitation goals of the stroke survivor (see Table 3). Stroke survivors with moderate to good hand function, who are likely to possess some range of motion in the hand, can grip, grasp, or pinch [30,31] hand rehabilitation devices and so have upper limb rehabilitation goals aimed at strengthening the existing motor ability. These goals may be related to improving grip strength and endurance, the ability to release objects or release time, the existing range of upper limb movements, and finger dexterity and regaining the ability to manipulate small objects (see Table 3). However, those with poor hand function, especially those with an ARAT score of 0 who cannot grasp, grip, or pinch objects irrespective of the sizes [31], have upper limb rehabilitation goals that focus on recovering some movement in the joints (shoulder, elbow, wrist, and/or fingers; see Tables 3 and 4).

Muscle weakness and the appearance of muscle stiffness, tightness, or tone (evident by the presence of a clenched hand) were clinically examined as being responsible for the poor hand function of the participants in this study (see Figure 3). The appearance of clenched hands has been reported as a clinical feature of spasticity [32]; moreover, the presence of muscle stiffness, tightness, and tone have all been connected with spasticity [33,34]. Previous studies have reported both spasticity and muscle weakness as the 2 major motor impairments following a stroke [35,36]. The severity of these impairments led to difficulty in hand immobility in 80% of those with poor hand function (with an ARAT score of 0), and according to an
earlier report [36], spasticity and muscle weakness can result in immobility.

Figure 3. Participants with poor hand function taking part in the 8-week rehabilitation exercise.

**Strengths**

The *UK National Clinical Guideline for Stroke* stipulates that stroke survivors should be considered for rehabilitation at any point after the stroke to potentially gain benefits [37]. However, an earlier study [38] that measured the accuracy of physical therapists’ early prediction of upper limb function reported that stroke survivors with ARAT scores more than 10 are those principally qualified to undergo rehabilitation exercises; this potentially excludes stroke survivors with poor hand function from taking part in hand rehabilitation. This study shows that not all stroke survivors with poor hand function should be considered ineligible to make use of hand rehabilitation devices, as those with some range of motion in their hand, as seen in participants with ARAT scores of 7 and 9 (see Table 4), can still benefit from hand rehabilitation devices and thus active hand rehabilitation.

**Limitations**

Only participants who exhibited poor hand function with an ARAT score of 0 were not able to benefit from active hand rehabilitation using devices. Those in this category whose poor hand function was due to muscle weakness were unable to carry out any intended active movement on the hand rehabilitation devices (see Table 4), even when supported to place their hand on them. In contrast, those whose poor hand function was due to hand stiffness or tightness, in addition to their inability to carry out intended active movement, were also faced with the problem of accessibility, which made it difficult for them to fit the device.

A limitation of this study was the inability to assess these conditions (muscle weakness and muscle tone or tightness)—examined to be responsible for the poor hand function—using the relevant outcome measures, such as motricity index, grip strength or pinch strength (for muscle weakness), or the Modified Ashworth Scale (for spasticity) [39], to quantify their severity. However, their severity was such that the hand was not useful in carrying out any of the ARAT tasks [31], as indicated by an ARAT score of 0.

**Future Direction**

Improvement in technological advancement has led to the development of devices such as rehabilitation gloves (smart or robotic gloves) that can be useful in stretching the hands of stroke survivors with poor hand function without requiring their active participation [40,41]. However, only stroke survivors with low spasticity (who possess some range of active motion in the hand [42]) may be able to make use of these rehabilitation gloves [40]. This means those with considerable muscle stiffness resulting in difficulty in passive motion [42] are still unlikely to freely access these devices; thus, future design of rehabilitation devices for hand rehabilitation should consider the problem of device accessibility in people with poor hand function due to considerable muscle stiffness or tightness.

**Conclusions**

It is therefore concluded that not all stroke survivors with impairments in their hands can interact with the available hand rehabilitation technologies, as those with an ARAT score of 0 cannot actively interact with any hand rehabilitation device. Thus, the selection of devices for hand rehabilitation should first consider the hand function of the affected stroke survivor. Since muscle stiffness or tightness in the hand results in poor hand function that can impede access to hand rehabilitation devices, future design of devices for hand rehabilitation should consider the accessibility needs of those with poor hand function as a result of hand stiffness or tightness. A similar observational study involving more stroke survivors will help ascertain the percentage of stroke survivors who fall into the category of having poor hand function and is therefore recommended.

**Acknowledgments**

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Data Availability
The data sets generated during this study are available from the corresponding author upon reasonable request.

Authors' Contributions
All authors contributed to the study’s methodology, investigation, and administration. Specifically, COW was involved with the conceptualizing, original draft writing, formal analysis, and visualization of the work. GS and MS were involved with editing and review of the draft, and AK was involved with the supervision of the project.

Conflicts of Interest
None declared.

References
18. Myers M. Rehab device enables stroke survivors with arm disabilities to do more training. Imperial College London. 2021 Sep 1. URL: https://tinyurl.com/wxydv9dc [accessed 2022-02-11]


Abbreviations

**ARAT**: Action Research Arm Test