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The impact of COVID-19 on the digital divide: A rapid review

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The impact of COVID-19 on the digital divide: A rapid review

3500 words

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Keywords

Digital divide; Health inequalities; COVID-19
Abstract

Objective

The increased reliance on digital technologies to deliver healthcare as a result of the COVID pandemic has meant pre-existing disparities in digital access and utilisation of healthcare might be exacerbated in disadvantaged patient populations. The aim of this rapid review conducted in January 2021 was to identify how this ‘digital divide’ was manifest during the first wave of the pandemic and highlight any areas which might be usefully addressed for the remainder of the pandemic and beyond.

Design

Rapid review and narrative synthesis

Data sources

The major medical databases including PubMed and Embase were searched alongside a hand-search of bibliographies.

Eligibility criteria

Original research papers available in English which described studies conducted since the beginning of the COVID-19 pandemic spring 2020.

Results

The search was described using PRISMA and identified nine studies. The results are presented within a novel framework consisting of the three key domains of the digital divide; 1) Digital Access, where a study described continuing issues with internet connectivity amongst vulnerable patients in the UK where described. 2) Digital Literacy, where seven studies described how ethnic minorities and the elderly were less likely to use digital technologies in accessing care. 3) Digital Assimilation,
where one study described how video technologies can reduce feelings of isolation and another how elderly black males were the most likely group to share information about COVID-19 on social media platforms.

Conclusions

Familiar difficulties in utilisation of digital healthcare amongst the elderly and ethnic minorities continue to be observed despite the unprecedented risk associated with direct contact with care providers. This is a further reminder that the digital divide is a persistent challenge that needs to be urgently addressed when considering the likelihood that in many instances these digital technologies are likely to remain at the centre of healthcare delivery.

Strengths and limitations of this study

The longstanding societal variations in the access and utilisation of digital technologies (A.K.A. the digital divide) in the pursuit of healthcare is widely recognised.

The enforced reliance on digital healthcare as a result of the COVID pandemic is likely to exacerbate the digital divide and further impact disadvantaged groups.

Our rapid review provided evidence of how pre-existing societal disparities in the access and utilisation of health-related digital technologies were accentuated by COVID-19.

The findings were presented within a novel framework that provides a comprehensive context for this and future explorations of the digital divide.
Introduction

A growing range of digital tools have been developed to help patients track their condition, connect with peer and clinical support, enable self-management, and aid more appropriate utilisation of health services (1, 2). When coordinated with the appropriate digital infrastructure they appear well placed to meet the need for more effective personalised healthcare, (3) that is capable of bridging the gap between increasing demand and restricted resource (4). The World Health Organisation’s (WHO) recently launched global strategy for digital health confirmed the expected role of digital technologies in creating a more equitable future for healthcare by offering “effective clinical and public health solutions to accelerate the achievement of the health and well-being... leaving none behind, [whether] children or adults, rural or urban.” (5).

Implicit within the digital transformation of healthcare and its role in reducing inequalities is that the relevant technologies are available across all levels of society (6, 7). However persistent discrepancies exist between communities in how they access and utilise digital technologies, differences compounded by the growing sophistication in the functionality of devices and connectivity (8). The result is that comparative advantages continue to be afforded to those groups that can maximise the capabilities of digital technologies (9, 10). These societal differences in access and adoption are commonly referred to as the “digital divide” (10), a catch-all phrase which implies a simple dichotomy but in reality describes a complex range of users whose level of adoption changes over time influenced by infrastructure, socio-economic environment and individual characteristics such as educational background and physical disability (11-15).

Despite acknowledged inequities in digital access and utilisation measures introduced to reduce infection rates following the onset of the COVID-19 pandemic in spring 2020. The understanding
that it was spread by person-to-person contact led to an acceleration of the reliance on digital health technologies both in Europe and the United States of America (6, 16-18). Because the spread of COVID was so rapid many of these digital interventions were introduced without the recommended periods of consultation and evaluation (19, 20). This led to concerns that the digital divide means these new models of healthcare delivery will disproportionately affect the health of disadvantaged communities (21-24) such as ethnic minorities (25), rural populations (26), the elderly (27) and residents of care homes (28). These concerns were heightened when it became apparent that the same groups on the “wrong” side of the digital divide were the most likely to experience severe symptoms and higher levels of mortality as a result of contracting the virus (21-24).

Many of these novel digitally-reliant processes that in March 2020 were considered a short-term fix are now becoming embedded in existing systems of care in the United Kingdom and elsewhere (17, 29). Therefore it is important to understand the implications of these new systems for patients and the quality and safety of the care they receive. This rapid review aims to explore how the digital divide manifested during the first wave of COVID-19 generating knowledge that can improve digital inclusion for the remainder of the pandemic and beyond.

**Methods**

**Study design**

Rapid reviews have previously been recommended for their ability to provide timely and credible evidence (30) and this review will draw on the major medical databases using the principles of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) to describe the
search (31). The results are presented within a novel framework that enables a more systematic
description of the various aspects of the digital divide explored by each study (see Box 1).

**Search strategy**

We searched PubMed, Embase, and Google Scholar, alongside hand-searches of bibliographies to
identify relevant manuscripts. In doing so we used a combination of the search terms ‘COVID 19’ or
‘pandemic’ or ‘COVID’ and ‘digital health’ or ‘telemedicine’ or ‘remote access’ or ‘digital divide’ to
identify studies which had explored the access or utilisation of information or communication
technologies in relation to health and care since the onset of the COVID-19 pandemic.

**Inclusion criteria**

To be included in our review, the manuscript must consist of original research specific to individuals
using digital technologies in relation to their health or well-being since the beginning of the
pandemic recognised by the WHO as March 2020 until January 2021 (32). This includes the
diagnosis, monitoring, or treatment of COVID-19 and any other condition or disease. The focus of
our work was the provision of care within the developed world to ensure relevance for
policymakers, commissioners and providers in these areas and so we limited the papers included to
those that were available in English.

**Study selection**

The process followed the four-stages of PRISMA (31); identification, screening, eligibility and final
inclusion and the search data presented in the PRISMA diagram (see Figure 1). This involved two
reviewers (IL and SG) screening titles, abstracts and where appropriate full texts against the
inclusion criteria and the final selection of papers agreed by both.

Figure 1 PRISMA diagram

Analysis procedures

We developed a framework that built on the work of Ai-chi Loh et al (33) to reflect a more nuanced
representation of the digital divide by describing it within three key domains; Digital Access relating
to the ability to access devices and internet; Digital Literacy describing the skill set of individuals,
and Digital Assimilation addressing the degree to which digital technologies are incorporated into
everyday life. Each domain consists of a series of related constructs and these are further defined
and presented with examples of each in Box 1. A descriptive summary of the characteristics of
each included study was produced (see Table 1) and the findings from the identified papers are
analysed within each of the key domains of the framework.

Patient and public involvement

No patients or members of the public were involved in the conceptualisation, design or
undertaking of this rapid review.
## Box 1 Framework for interpreting the digital divide in healthcare (after Ai-chi Loh et al (33))

<table>
<thead>
<tr>
<th>Domain</th>
<th>Definition</th>
<th>Construct</th>
<th>Definition</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Digital Access</td>
<td>The ability to access the necessary hardware, software, and internet services associated with utilisation of digital technologies (34).</td>
<td>The types of device available</td>
<td>The nature and functionality of the digital device (10).</td>
<td>The model of smartphone or desktop computer and any peripheral or supporting technology such as hard drives or printers.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>The ease with which devices can be accessed</td>
<td>How readily individuals can access digital devices (8, 10).</td>
<td>Relying on the local library for access to a computer.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>The autonomy and reliability of internet connectivity</td>
<td>The degree of independence with which the internet can be reliably accessed (35, 36).</td>
<td>Consistent access to the internet via a user’s home internet network.</td>
</tr>
<tr>
<td>Digital Literacy</td>
<td>The degree of sophistication with which individuals are able to use digital technologies (37).</td>
<td>Digital skill set</td>
<td>The confidence and ability of an individual to utilise a variety of digital technologies (38).</td>
<td>The ability to use and manage email.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Types of digital usage</td>
<td>The ways in which digital technologies are utilised (39).</td>
<td>Using search engines to access information on current affairs</td>
</tr>
<tr>
<td>Digital Assimilation</td>
<td>The degree to which digital technologies are incorporated and utilised in everyday life (34, 40).</td>
<td>Engagement with digital technologies</td>
<td>The degree to which individuals use digital technologies to enhance social connections and values (41).</td>
<td>Establishing a community group on a social media platform to support elderly neighbours.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Social support</td>
<td>Social connections that facilitate an individual's engagement with digital technologies (38).</td>
<td>The availability of technical support in the use of digital technologies from a son or daughter.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Harnessing digital outcomes</td>
<td>The ability to contextualise the use of digital technologies to achieve quantifiable outputs (42, 43).</td>
<td>Using software apps to reach and maintain fitness goals.</td>
</tr>
</tbody>
</table>
Results

A total of 28 candidate articles were identified from a search of the named databases and hand-searches from the bibliographies of these references. Ultimately nine papers were selected for the analysis, the remaining papers were excluded as they were either opinion pieces that did not contain original research or despite being published after March 2020 referenced research conducted prior to the onset of the epidemic.

One study looked at Digital Access; set in UK primary care it explored internet connectivity amongst vulnerable patients (44). It was also one of the seven studies that looked at Digital Literacy (44) alongside five studies set in the United States that explored the use of digital technologies in accessing care amongst different ethical groups (45-49) and one study conducted in Italy that looked at the age and gender of patients using telemedicine (50). Two studies were concerned with Digital Assimilation, one set in Italy described the social support gained from using video messaging platforms (51) and a second again set in the United States explored the characteristics of individuals posting COVID-related content on social media (52). The key characteristics of these papers are summarised in Table 1.
<table>
<thead>
<tr>
<th>Authors</th>
<th>Study design</th>
<th>Country</th>
<th>Study Population</th>
<th>Research question</th>
<th>Analytical framework</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Campos-Castillo &amp;, Anthony</td>
<td>Cohort Study</td>
<td>USA</td>
<td>10,624 USA-wide survey respondents</td>
<td>What are the characteristics of patients that used ICTs to connect with care providers in relation to COVID-19.</td>
<td>Logistic regression</td>
<td>Total of 17% of respondents self-reported using telehealth because of the pandemic. Black respondents were more likely than Whites to report using telehealth because of the pandemic (OR, 1.42; 95% CI, 1.07–1.88).</td>
</tr>
<tr>
<td>Castillo &amp;, Anthony (2020)</td>
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<td></td>
</tr>
<tr>
<td>Campos-Castillo C &amp;, Laestadius</td>
<td>Cohort study</td>
<td>USA</td>
<td>10,541 USA-wide survey respondents</td>
<td>What are the characteristics of patients posting COVID-19 related messages on social media</td>
<td>Logistic regression</td>
<td>Respondents who identified as black (OR 1.29, 95% CI 1.02-1.64; P=.03), or Latino (OR 1.66, 95% CI 1.36-2.04; P=.03) had higher odds than respondents who identified as white of reporting that they posted COVID-19 content on social media.</td>
</tr>
<tr>
<td>Chunara R, et al (2020)</td>
<td>Cohort study</td>
<td>USA</td>
<td>140,184 patients</td>
<td>What are the characteristics of patients that use telemedicine</td>
<td>Descriptive statistics and logistic regression</td>
<td>Black patients nearly half as likely as white patients to access care through telemedicine (OR 0.6 times (95% CI: 0.58–0.63).</td>
</tr>
<tr>
<td>Eberly, et al. (2020)</td>
<td>Cohort study</td>
<td>USA</td>
<td>2,940 cardiovascular outpatients</td>
<td>What are the characteristics of patients that complete a telemedicine consultation</td>
<td>Logistic regression</td>
<td>Being female and being non-English speaking were independently associated with less telemedicine use.</td>
</tr>
<tr>
<td>Gabbiadini et al 2020 (51)</td>
<td>Cross-sectional</td>
<td>Italy</td>
<td>465 respondents</td>
<td>Whether the use ICT promoted perceptions of social support (mitigating the psychological effects of lockdown)</td>
<td>Separate multiple and simple regression models</td>
<td>The amount of technology use was a significant predictor of perceived social support. (OR 2.40, p &lt; 0.02, 99% CI –0.01, 0.51).</td>
</tr>
<tr>
<td>Study</td>
<td>Study Type</td>
<td>Country</td>
<td>Description</td>
<td>Methodology</td>
<td>Findings</td>
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<tr>
<td>Hughes et al</td>
<td>Mixed methods</td>
<td>UK</td>
<td>156 high-risk individuals and a further 1,217 vulnerable patients over the age of 70</td>
<td>Can medical students (general practitioner trainees) use teleconsultations to assess the needs of patients and support digital access to healthcare</td>
<td>A total of 22% high risk patients and 44% of vulnerable patients reported connectivity issues. Participants reported that were confident in ordering medication online.</td>
<td></td>
</tr>
<tr>
<td>Runfola et al</td>
<td>Cohort study</td>
<td>Italy</td>
<td>33 Bariatric outpatients</td>
<td>What are the characteristics of patients that completed a telemedicine consultation</td>
<td>Categorical data was compared using the chi-square test. Continuous variables compared using the Student’s t test.</td>
<td></td>
</tr>
<tr>
<td>Weber et al</td>
<td>Cohort study</td>
<td>USA</td>
<td>52,585 patients diagnosed with COVID-19</td>
<td>What are the characteristics of patients that access care by telemedicine, ER or office visit</td>
<td>Descriptive statistics and multinomial regression analysis. White patients had the highest predicted probabilities of using telehealth (46.7%) Black and Hispanic patients over 65 have lowest predicted probability (11.3%)</td>
<td></td>
</tr>
</tbody>
</table>
| Whaley et al     | Cross-sectional     | USA      | Data from 5.6 to 6.8 million US individuals with employer health insurance between 2018, and 2020, | What are the characteristics of patients that use telemedicine. | Logistic regression. Patients living in post codes with lower-income or majority racial/ethnic minority populations had lower rates of adoption of telemedicine; 
≥80% racial/ethnic minority post code: -71.6 per 10,000 (95% CI, -87.6 to -55.5); 
79%-21% racial/ethnic minority post code: -15.1 per 10,000 (95% CI, -19.8 to -10.4). |
Digital Access

We identified one study concerned with the access of digital technologies, specifically the reliability of internet connectivity. It was conducted in United Kingdom (UK) primary care as part of a study whose overall aim was to explore whether vulnerable patients might be usefully supported by tele-coaching in the use of digital health technologies, in this instance by General Practitioner (GP) trainees (44). As part of these conversations a direct question was asked around internet connectivity and the authors reported that 22% of high-risk patients and 44% of vulnerable patients reported issues (44).

Digital Literacy

A total of seven studies addressed the domain of Digital Literacy and in particular an individual’s digital skill set, specifically in relation to the ways in which they accessed care. All provided comparisons of use between groups using descriptions of demographic characteristics that included age, gender and ethnicity (53). Two studies used routinely collected electronic health data, though were conducted independently of each other, at two different sites in New York (United States); Chunura et al (2020) used data gathered from patients at New York University Hospital collected over a 6-week period to determine whether they had received their COVID-19 diagnosis at an office visit or via video consultation. The authors described that the digital infrastructure of the service was well resourced and established and therefore attributed the reduced utilization of telemedicine by black patients to factors unrelated to the digital capacity of the facility (45). The second study set in New York was also situated within a large health care centre and again compared the means of accessing healthcare between ethnic groups within the early months of the pandemic (46). They found that black and Hispanic patients were more likely to visit the Emergency Room or arrange an office visit than use telehealth than their white or Asian...
counterparts (46). In this instance the authors recognised that the more extensive use of ER may be
due to the disproportionate number of ethnic minorities that experienced severe COVID-related
symptoms (46). Another study set in the United States compared the use of telemedicine amongst
commercially insured patients from 2018 through to 2020 (47). In doing so they explored
differences in both the nature of the care they received and the means of access in the first two
months of the pandemic and described that though there was an increase in telemedicine it did not
make up the shortfall in the number of visits in comparison to the usual levels of assessing
preventative or elective care amongst ethnic minorities (47). Campos-Castilo and Anthony
carried out a secondary analysis of a national survey a secondary analysis of cross-sectional survey
data from the Pew Research Center’s American Trends Panel, which is a national, probability-based
online panel of adults (18 or older) living in U.S. households exploring self-reported use of
telemedicine and following adjustment for socio-economic status (SES), age and perceived level of
threat to their health from the pandemic (no threat, minor, or major) (49). They found black
patients were actually more likely to contact care providers using ICTs if they perceived their health
was threatened by the virus (49).

Two studies specifically explored whether there were differences in the characteristics of patients
fulfilling pre-arranged or routine video-consultations during the pandemic. One of these studies
was also set in the USA and compared the characteristics of cardiovascular patients that ‘attended’
teleconsultations and found no differences in cancellation rates based on race, ethnicity, or
household income. However, differences between genders were observed with those completing
telemedicine tending to be male and older (48). In Italy Runfola et al explored the utilisation and
subsequent satisfaction with video consultations amongst a group of bariatric patients. They found
no significant differences in terms age or gender between those that succeeded or failed to
complete a video call (50). However, in terms of overall numbers just under 58% of patients
fulfilled the video-consultation and the authors felt that this was due to the absence of basic computer skills and a lack of self-efficacy in utilising video-call systems (50). In relation to self-efficacy the Hughes study set in the UK also assessed vulnerable patients’ confidence and ability to order medications online and reported they were comfortable and confident with the process (44).

**Digital Assimilation**

Two studies explored the use of digital technologies in relation to maintaining or interacting with a social network. One study set in Italy described how feelings of loneliness, boredom and irritability were all reduced as a result of regular utilisation of video calls, and the positive effects on maintaining meaningful relationships and mental health (51). Meanwhile in the United States another secondary analysis of the same cross-sectional survey data from the Pew Research Center’s American Trends Panel, was conducted to understand if there were differences in the characteristics of individuals that posted Covid-19 related content to social media platforms (52). The authors discovered that proportionally members of racial and ethnic minority groups and amongst these older black males, were the most likely to contribute Covid-19 related content (52).
Discussion

General findings

Our rapid review identified how pre-existing societal disparities in access to and utilisation of health-related digital technologies were accentuated by COVID-19. We identified nine studies that explored various constructs within the three domains of our novel digital divide framework. In relation to Digital Access, poor internet access amongst the elderly was reported (44), as regards Digital Literacy lower levels of take-up of telemedicine amongst certain communities in the United States were described particularly among black and Hispanic patients (46-49). Within the domain of Digital Assimilation one study described how face-time technology can sustain relationships amongst dislocated peer groups (50), and another how black and elderly males, previously considered a group unprepared to share health information on social media platforms were the demographic most likely to post content on the pandemic, an important consideration in understanding the emerging scepticism of the COVID vaccine in ethnic groups (51).

That our search uncovered so few studies can be attributed to two factors relating to the pandemic; first that the research capacity of healthcare organisations would have been compromised by dealing with the exceptional demand on their services (54, 55); second that the issue of the ‘digital divide’ which had previously failed to be considered a priority was unlikely to be addressed during the most serious public health crisis in a generation (56).

Strengths and weaknesses

Although our search initially uncovered numerous titles many were opinion or editorial pieces, demonstrating how widely recognised the phenomenon of the digital divide is but also its lack of
priority as a subject for original research (22, 26, 57-59). Though the studies identified were conducted within only three countries to date they constituted three of the top four worst death rates per capita from COVID-19 (60, 61). All offered valuable insight into how the digital divide was reflected in the early stages of the pandemic though the lack of a theoretical underpinning limited generalisability (61) and that two of the studies relied on self-reported data (44, 49) raises familiar issues regards their reliability (62).

Specific findings

Digital Access

The Hughes paper provides the latest example of how discrepancies in reliable internet connectivity continue in England (44, 63) findings which were corroborated by the most recent surveys of digital access conducted in the UK which found that nearly 7% of homes in England and Wales did not have a reliable internet connection (64, 65), a lack of connectivity that disproportionately affected the elderly, those of lower socio-economic status, and the disabled (64-67).

Despite the calls to harness digital technologies on a global scale (5, 68), these also need to address the stubborn differences in digital access that remain within the developed world where significant divisions in digital connectivity and utility remain and continue to affect the most vulnerable members of society (8, 68-77). The pandemic prompted broader acknowledgement of these differences in several health economies where a number of initiatives were introduced (59, 78). For example in the UK broadband providers lowered prices and reduced data caps for the vulnerable (79), and in the United States roving buses were used to provide Wi-Fi access for unconnected communities (80).
Digital Literacy

The patterns in digital literacy relating to socio-economic status, age, or race described in four of the studies we identified (45, 46, 48, 49) have been observed for nearly three decades (8, 16-18, 66, 67, 81-83). However, prior to the pandemic, using traditional methods of in-person access did not hold the same degree of risk as during a pandemic where airborne transmission led to widespread recommendations to minimise social contact (32). This may be due in part to variations in individual perception of risk influenced by personal experience, social values, and the attitudes of friends and family (84). It also reflects the resistance of the digital divide to intervention. A number of previous attempts have been made to connect less technologically enabled patients to the appropriate care (58, 85, 86). However, the non-adoption and abandonment of telehealth technologies by the intended users is common (87-90) complicated by the influences of the provider organisation and the design and compatibility of the intervention (91). Self-efficacy, patient activation and motivation are also critical yet underexplored components of the uptake of digital technology (92) as are the impact of patients’ knowledge of their condition; the expectations of the care they should receive, their social situation and the resources at their disposal (93).

In attempting to unpick this complexity a number of theoretical frameworks have been developed intended to support adoption and produce transferable learning for a range of digital innovations (94). There have also been calls for greater patient and public involvement in designing and developing digital healthcare to ensure the needs and preferences of the full range of patients are incorporated (95, 96).

Digital Assimilation
For over a decade the internet has been recognised as a key source of health information for the public and patients, yet the precise role of social media in the communication of health-related information is less clear (97). Although limited, evidence tended to suggest that sharing health information online was favoured by the young (98) and was less so amongst the elderly or those of lower SES (99). However, one study we found described how older black males were more likely to share information about COVID through social media channels than other demographic groups (52). This may in part be due to the growing reluctance amongst black and ethnic minority groups to trust information provided by healthcare professionals or the mainstream media (100-102). That highlights how the growing consumption of health information through a largely unregulated network of social media platforms can have serious repercussions for public health (102-106). This is of particular concern when placed in the context of the growing scepticism about the COVID-19 vaccine in minority communities (107, 108).

Despite the potential for spreading misinformation the work by Runfola observed benefits for mental health from the use of face-to-face digital contact during the pandemic (50). The last five years has seen a growing realisation that the responsible use of social media can be an effective means of alleviating depression and social isolation and improve mental well-being (109-111). In particular the utilisation of face-time technologies has been shown to increase and enhance social interactions (112) and engagement (113). During the pandemic these benefits were recognised by the UK government in their scheme that provided free tablets to care homes to help connect isolating residents with their families and loved ones (114).

Conclusions
The rapid incorporation of digital technologies into mainstream health care delivery due to the COVID pandemic was widely understood and accepted by patients unwilling to breach social distancing advice. However, not all patient groups were either willing or able to utilise the digital services made available nor maximise the reported benefits of face-time technology to alleviate the effects of isolation. Our findings provide further evidence that patient engagement with any model of digital healthcare is vulnerable to complex socio-political factors. If more are to reap the potential benefits of digital healthcare then not only are acknowledged improvements in infrastructure needed but also concerted efforts to train, equip and motivate all patients in its use.

**Ethics statement**

Ethical approval was not required

**Data Sharing Statement**

No additional data is available for sharing.

**Funding**

The work was not externally funded

**Conflicts of interest**

There are no conflicts of interest

**Author’s contribution**

IL was responsible for the conception of the work and the design of the review. IL and SG reviewed the articles and IL led the drafting of the article with input from SG and DS. SG and DS all provided critical revisions. The final version was drafted by IL and approved by SG and DS.
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Reference List

4. Rimmer A. Technology will improve doctors’ relationships with patients, says Topol review. BMJ. 2019;364:l661.
8. van Deursen AJAM, van Dijk JAGM. The first-level digital divide shifts from inequalities in physical access to inequalities in material access. New Media & Society. 2018;21(2):354-75.


64. Sweaney M. Slow digital services are marginalising rural areas, MPs warn. The Guardian. 2019.

108. Jones Z. Why some black americans are sceptical of the covid-19 vaccine 2020 [ ]

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Figure I PRISMA diagram

- Records identified through database searching (n = 25)
- Additional records identified through other sources (n = 3)

Records after duplicates removed (n = 28)

Records screened (n = 28)

- Records excluded (n = 18)
- Full-text articles assessed for eligibility (n = 9)

- Full-text articles excluded, as concerned with broadband access only (n = 1)

Studies included in narrative synthesis (n = 9)
<table>
<thead>
<tr>
<th>Section and Topic</th>
<th>Item #</th>
<th>Checklist item</th>
</tr>
</thead>
<tbody>
<tr>
<td>TITLE</td>
<td>1</td>
<td>Identify the report as a systematic review.</td>
</tr>
<tr>
<td>ABSTRACT</td>
<td>2</td>
<td>See the PRISMA 2020 for Abstracts checklist.</td>
</tr>
<tr>
<td>INTRODUCTION</td>
<td>3</td>
<td>Describe the rationale for the review in the context of existing knowledge.</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>Provide an explicit statement of the objective(s) or question(s) the review addresses.</td>
</tr>
<tr>
<td>METHODS</td>
<td>5</td>
<td>Specify the inclusion and exclusion criteria for the review and how studies were grouped for the syntheses.</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>Specify all databases, registers, websites, organisations, reference lists and other sources searched or consulted to identify studies. Specify the date when each source was last searched or consulted.</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>Present the full search strategies for all databases, registers and websites, including any filters and limits used.</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>Specify the methods used to decide whether a study met the inclusion criteria of the review, including how many reviewers screened each record and each report retrieved, whether they worked independently, and if applicable, details of automation tools used in the process.</td>
</tr>
<tr>
<td></td>
<td>9</td>
<td>Specify the methods used to collect data from reports, including how many reviewers collected data from each report, whether they worked independently, any processes for obtaining or confirming data from study investigators, and if applicable, details of automation tools used in the process.</td>
</tr>
<tr>
<td></td>
<td>10a</td>
<td>List and define all outcomes for which data were sought. Specify whether all results that were compatible with each outcome domain in each study were sought (e.g. for all measures, time points, analyses), and if not, the methods used to decide which results to collect.</td>
</tr>
<tr>
<td></td>
<td>10b</td>
<td>List and define all other variables for which data were sought (e.g. participant and intervention characteristics, funding sources). Describe any assumptions made about any missing or unclear information.</td>
</tr>
<tr>
<td></td>
<td>11</td>
<td>Specify the methods used to assess risk of bias in the included studies, including details of the tool(s) used, how many reviewers assessed each study and whether they worked independently, and if applicable, details of automation tools used in the process.</td>
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<tr>
<td></td>
<td>12</td>
<td>Specify for each outcome the effect measure(s) (e.g. risk ratio, mean difference) used in the synthesis or presentation of results.</td>
</tr>
<tr>
<td>Synthesis methods</td>
<td>13a</td>
<td>Describe the processes used to decide which studies were eligible for each synthesis (e.g. tabulating the study intervention characteristics and comparing against the planned groups for each synthesis (item #5)).</td>
</tr>
<tr>
<td></td>
<td>13b</td>
<td>Describe any methods required to prepare the data for presentation or synthesis, such as handling of missing summary statistics, or data conversions.</td>
</tr>
<tr>
<td></td>
<td>13c</td>
<td>Describe any methods used to tabulate or visually display results of individual studies and syntheses.</td>
</tr>
<tr>
<td></td>
<td>13d</td>
<td>Describe any methods used to synthesize results and provide a rationale for the choice(s). If meta-analysis was performed, describe the model(s), method(s) to identify the presence and extent of statistical heterogeneity, and software package(s) used.</td>
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<td></td>
<td>13e</td>
<td>Describe any methods used to explore possible causes of heterogeneity among study results (e.g. subgroup analyses, meta-regression).</td>
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<td></td>
<td>13f</td>
<td>Describe any sensitivity analyses conducted to assess robustness of the synthesized results.</td>
</tr>
<tr>
<td>Reporting bias assessment</td>
<td>14</td>
<td>Describe any methods used to assess risk of bias due to missing results in a synthesis (arising from reporting bias).</td>
</tr>
<tr>
<td>Certainty assessment</td>
<td>15</td>
<td>Describe any methods used to assess certainty (or confidence) in the body of evidence for an outcome.</td>
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<tr>
<td>Section and Topic</td>
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<tr>
<td>Study selection</td>
<td>16a</td>
<td>Describe the results of the search and selection process, from the number of records identified in the search to the number of studies included in the review, ideally using a flow diagram.</td>
</tr>
<tr>
<td></td>
<td>16b</td>
<td>Cite studies that might appear to meet the inclusion criteria, but which were excluded, and explain why they were excluded.</td>
</tr>
<tr>
<td>Study characteristics</td>
<td>17</td>
<td>Cite each included study and present its characteristics.</td>
</tr>
<tr>
<td>Risk of bias in studies</td>
<td>18</td>
<td>Present assessments of risk of bias for each included study.</td>
</tr>
<tr>
<td>Results of individual studies</td>
<td>19</td>
<td>For all outcomes, present, for each study: (a) summary statistics for each group (where appropriate) and (b) an effect estimate and its precision (e.g. confidence/credible interval), ideally using structured tables or plots.</td>
</tr>
<tr>
<td>Results of syntheses</td>
<td>20a</td>
<td>For each synthesis, briefly summarise the characteristics and risk of bias among contributing studies.</td>
</tr>
<tr>
<td></td>
<td>20b</td>
<td>Present results of all statistical syntheses conducted. If meta-analysis was done, present for each the summary estimate and its precision (e.g. confidence/credible interval) and measures of statistical heterogeneity. If comparing groups, describe the direction of the effect.</td>
</tr>
<tr>
<td></td>
<td>20c</td>
<td>Present results of all investigations of possible causes of heterogeneity among study results.</td>
</tr>
<tr>
<td></td>
<td>20d</td>
<td>Present results of all sensitivity analyses conducted to assess the robustness of the synthesized results.</td>
</tr>
<tr>
<td>Reporting biases</td>
<td>21</td>
<td>Present assessments of risk of bias due to missing results (arising from reporting biases) for each synthesis assessed.</td>
</tr>
<tr>
<td>Certainty of evidence</td>
<td>22</td>
<td>Present assessments of certainty (or confidence) in the body of evidence for each outcome assessed.</td>
</tr>
<tr>
<td>DISCUSSION</td>
<td>23a</td>
<td>Provide a general interpretation of the results in the context of other evidence.</td>
</tr>
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<td></td>
<td>23b</td>
<td>Discuss any limitations of the evidence included in the review.</td>
</tr>
<tr>
<td></td>
<td>23c</td>
<td>Discuss any limitations of the review processes used.</td>
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<td></td>
<td>23d</td>
<td>Discuss implications of the results for practice, policy, and future research.</td>
</tr>
<tr>
<td>OTHER INFORMATION</td>
<td>24a</td>
<td>Provide registration information for the review, including register name and registration number, or state that the review was not registered.</td>
</tr>
<tr>
<td>Registration and protocol</td>
<td>24b</td>
<td>Indicate where the review protocol can be accessed, or state that a protocol was not prepared.</td>
</tr>
<tr>
<td></td>
<td>24c</td>
<td>Describe and explain any amendments to information provided at registration or in the protocol.</td>
</tr>
<tr>
<td>Support</td>
<td>25</td>
<td>Describe sources of financial or non-financial support for the review, and the role of the funders or sponsors in the review.</td>
</tr>
<tr>
<td>Competing interests</td>
<td>26</td>
<td>Declare any competing interests of review authors.</td>
</tr>
<tr>
<td>Availability of data, code and other materials</td>
<td>27</td>
<td>Report which of the following are publicly available and where they can be found: template data collection forms; data extracted from included studies; data used for all analyses; analytic code; any other materials used in the review.</td>
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# The impact of COVID-19 on the digital divide: A rapid review

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<td>Date Submitted by the Author:</td>
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| Complete List of Authors: | Litchfield, Ian; University of Birmingham College of Medical and Dental Sciences, Institute of Applied Health Research  
Shukla, David; University of Birmingham College of Medical and Dental Sciences, Institute of Applied Health Research  
Greenfield, Sheila; University of Birmingham College of Medical and Dental Sciences, Institute of Applied Health Research |
| Primary Subject Heading: | Health services research |
| Secondary Subject Heading: | Communication, Global health, Health policy, Patient-centred medicine |
| Keywords:       | COVID-19, Organisation of health services < HEALTH SERVICES ADMINISTRATION & MANAGEMENT, Quality in health care < HEALTH SERVICES ADMINISTRATION & MANAGEMENT |
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The impact of COVID-19 on the digital divide: A rapid review

3500 words

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Keywords

Digital divide; Health inequalities; COVID-19
Abstract

Objective

The increased reliance on digital technologies to deliver healthcare as a result of the COVID-19 pandemic has meant pre-existing disparities in digital access and utilisation of healthcare might be exacerbated in disadvantaged patient populations. The aim of this rapid review was to identify how this ‘digital divide’ was manifest during the first wave of the pandemic and highlight any areas which might be usefully addressed for the remainder of the pandemic and beyond.

Design

Rapid review and narrative synthesis

Data sources

The major medical databases including PubMed and Embase and Google Scholar were searched alongside a hand-search of bibliographies.

Eligibility criteria

Original research papers available in English which described studies conducted since the 1st of March 2020 until 31st July 2021.

Results

The search was described using PRISMA and identified nine studies. The results are presented within a refined framework describing the three key domains of the digital divide; 1) Digital Access, within which one study described continuing issues with internet connectivity amongst vulnerable patients in the UK. 2) Digital Literacy, where seven studies described how ethnic minorities and the elderly were less likely to use digital technologies in accessing care. 3) Digital Assimilation, where one study described how video technologies can reduce feelings of isolation and another how
elderly black males were the most likely group to share information about COVID-19 on social media platforms.

Conclusions

During the early phase of the pandemic in the developed world, familiar difficulties in utilisation of digital healthcare amongst the elderly and ethnic minorities continued to be observed. This is a further reminder that the digital divide is a persistent challenge that needs to be urgently addressed when considering the likelihood that in many instances these digital technologies are likely to remain at the centre of healthcare delivery.

Strengths and limitations of this study

This rapid review provides timely information on the impact of COVID-19 on the digital divide during the first wave of the pandemic.

The findings were presented within a framework developed to provide a more comprehensive context for this and future explorations of the digital divide.

The search covered three key databases and was augmented by manual searches though the quality of the papers identified was not formally assessed.
Introduction

A growing range of digital tools have been developed to help patients track their condition, connect with peer and clinical support, enable self-management, and aid more appropriate utilisation of health services (1, 2). When coordinated with the appropriate digital infrastructure they appear well placed to meet the need for more effective personalised healthcare, (3) that is capable of bridging the gap between increasing demand and restricted resource (4). The World Health Organisation's (WHO) recently launched global strategy for digital health confirmed the expected role of digital technologies in creating a more equitable future for healthcare by offering "effective clinical and public health solutions to accelerate the achievement of the health and well-being... leaving none behind, [whether] children or adults, rural or urban.” (5).

Implicit within the digital transformation of healthcare and its role in reducing inequalities is that the relevant technologies are available across all levels of society (6, 7). However persistent discrepancies exist across geographies and between communities in how they access and utilise digital technologies, differences compounded by the growing sophistication in the functionality of devices and connectivity (8-10). The result is that comparative advantages continue to be afforded to those groups that can maximise the capabilities of digital technologies (11, 12). These societal differences in access and adoption are commonly referred to as the “digital divide” (12), a catch-all phrase which implies a simple dichotomy but in reality describes a complex range of users whose level of adoption changes over time influenced by infrastructure, socio-economic environment and individual characteristics such as educational background and physical disability (13-17).

Despite the acknowledged inequities in digital access and utilisation, measures introduced to reduce infection rates following the onset of the COVID-19 pandemic in spring 2020, led to an
acceleration of the reliance on digital health technologies both in Europe and the United States of America (6, 18-20). Because the spread of COVID-19 was so rapid many of these digital interventions were introduced without the recommended periods of consultation and evaluation (21, 22). This rapid introduction led to concerns that the new digitally reliant models of healthcare delivery will disproportionately affect the health of disadvantaged communities (23-26) such as ethnic minorities (27), rural populations (28), the elderly (29) and residents of care homes (30). These concerns were heightened when it became apparent that the same groups on the “wrong” side of the digital divide were the most likely to experience severe symptoms and higher levels of mortality as a result of contracting the virus (23-26).

Many of these novel digitally-reliant processes that in March 2020 were considered a short-term fix are now becoming embedded in existing systems of care in the United Kingdom and elsewhere (19, 31). Therefore it is important to understand the implications of these new systems for patients and the quality and safety of the care they receive. This rapid review aims to explore how the digital divide manifested during the first wave of COVID-19 generating knowledge that can improve digital inclusion for the remainder of the pandemic and beyond.

Methods

Study design

Rapid reviews have previously been recommended by the WHO amongst others for their ability to provide timely and credible evidence for policymakers and practitioners in what is a dynamic and evolving public health crisis (32)(33). We have used many of the principles of a systematic review process; our search terms were clearly defined using Boolean principles and the Preferred
Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) described the search (34). The inclusion and exclusion criteria were also clearly defined and two reviewers agreed on the selection of the various papers (The search terms can be found in supplementary file 1). However, the systematic review process was expedited by amending several steps i.e. drawing only on the major medical databases and forgoing a structured appraisal of the quality of selected studies in place of a transparent description of the characteristics of each within the results.

The results are presented within the three key domains of a framework (informed by Ai-Chi Loh et al’s work (35)) to enable a more systematic description of the various aspects of the digital divide explored by each study (see Box 1).

**Search strategy**

We searched PubMed, Embase, and Google Scholar, alongside hand-searches of bibliographies to identify relevant manuscripts. In doing so we used a combination of the search terms ‘COVID 19’ or ‘pandemic’ or ‘COVID’ and ‘digital health’ or ‘telemedicine’ or ‘remote access’ or ‘digital divide’ to identify studies which had explored the access or utilisation of information or communication technologies in relation to health and care since the onset of the COVID-19 pandemic. Bibliographies within the publications we identified were searched alongside a manual search.

**Inclusion criteria**

To be included in our review, the manuscript must consist of original research specific to individuals using digital technologies in relation to their health or well-being since the beginning of the pandemic recognised by the WHO as March 2020 with any publication published from the 1st of
March until the 31st of July 2021 (36). This includes the diagnosis, monitoring, or treatment of COVID-19 and any other condition or disease. The focus of our work was the provision of care within the developed world (i.e. one which is predominantly industrialized and more economically developed with a higher individual income (37) during the early phase of the pandemic to ensure relevance for policymakers, commissioners and providers in these areas and so we limited the papers included to those that were available in English.

Study selection

The process followed the four-stages of PRISMA (34); identification, screening, eligibility and final inclusion and the search data presented in the PRISMA diagram (see Figure 1). This involved two reviewers (IL and SG) screening titles, abstracts and where appropriate full texts against the inclusion criteria and the final selection of papers agreed by both.

Figure 1 PRISMA diagram

Analysis procedures

We developed a framework that built on the work of Ai-chi Loh et al (35) to reflect a more nuanced representation of the digital divide describing it within three key domains; Digital Access relating to the ability to access devices and internet; Digital Literacy describing the skill set of individuals, and Digital Assimilation addressing the degree to which digital technologies are incorporated into everyday life. Each domain consists of a series of related constructs and these are further defined and presented with examples of each in Box 1. A descriptive summary of the characteristics of each included study was produced (see Table 1) and the findings from the identified papers are analysed within each of the three domains of our refined framework.
Patient and public involvement

No patients or members of the public were involved in the conceptualisation, design or undertaking of this rapid review.
Box 1 Framework for interpreting the digital divide in healthcare (after Ai-chi Loh et al (35))

<table>
<thead>
<tr>
<th>Domain</th>
<th>Definition</th>
<th>Construct</th>
<th>Definition</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Digital Access</td>
<td>The ability to access the necessary hardware, software, and internet services associated with utilisation of digital technologies (38).</td>
<td>The types of device available</td>
<td>The nature and functionality of the digital device (12).</td>
<td>The model of smartphone or desktop computer and any peripheral or supporting technology such as hard drives or printers.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>The ease with which devices can be accessed</td>
<td>How readily individuals can access digital devices (8, 12).</td>
<td>Relying on the local library for access to a computer.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>The autonomy and reliability of internet connectivity</td>
<td>The degree of independence with which the internet can be reliably accessed (39, 40).</td>
<td>Consistent access to the internet via a user’s home internet network.</td>
</tr>
<tr>
<td>Digital Literacy</td>
<td>The degree of sophistication with which individuals are able to use digital technologies (41).</td>
<td>Digital skill set</td>
<td>The confidence and ability of an individual to utilise a variety of digital technologies (42).</td>
<td>The ability to use and manage email.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Types of digital usage</td>
<td>The ways in which digital technologies are utilised (43).</td>
<td>Using search engines to access information on current affairs.</td>
</tr>
<tr>
<td>Digital Assimilation</td>
<td>The degree to which digital technologies are incorporated and utilised in everyday life (38, 44).</td>
<td>Engagement with digital technologies</td>
<td>The degree to which individuals use digital technologies to enhance social connections and values (45).</td>
<td>Establishing a community group on a social media platform to support elderly neighbours.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Social support</td>
<td>Social connections that facilitate an individual’s engagement with digital technologies (42).</td>
<td>The availability of technical support in the use of digital technologies from a son or daughter.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Harnessing digital outcomes</td>
<td>The ability to contextualise the use of digital technologies to achieve quantifiable outputs (46, 47).</td>
<td>Using software apps to reach and maintain fitness goals.</td>
</tr>
</tbody>
</table>
Results

A total of 28 candidate articles were identified from a search of the named databases and hand-searches from the bibliographies of these references. Ultimately nine papers were selected for the analysis, the remaining papers were excluded as they were either opinion pieces that did not contain original research or despite being published after March 2020 referenced research conducted prior to the onset of the epidemic. One study looked at Digital Access; set in UK primary care it explored internet connectivity amongst vulnerable patients (including those that have received an organ transplant, are undertaking immunotherapy or an intense course of radiotherapy for lung cancer) (48, 49). It was also one of the seven studies that looked at Digital Literacy (49) alongside five studies set in the United States that explored the use of digital technologies in accessing care amongst different ethnic groups (50-54) and one study conducted in Italy that looked at the age and gender of patients using telemedicine (55). Two studies were concerned with Digital Assimilation, one set in Italy described the social support gained from using video messaging platforms (56) and a second again set in the United States explored the characteristics of individuals posting COVID-related content on social media (57). The key characteristics of these papers are summarised in Table 1.
Table 1 Summary of study characteristics

<table>
<thead>
<tr>
<th>Authors</th>
<th>Study design</th>
<th>Country</th>
<th>Study Population</th>
<th>Research question</th>
<th>Analytical framework</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Campos-Castillo &amp; Anthony (2020) (54)</td>
<td>Cohort Study</td>
<td>USA</td>
<td>10 624 USA-wide survey respondents</td>
<td>What are the characteristics of patients that used ICTs to connect with care providers in relation to COVID-19.</td>
<td>Logistic regression</td>
<td>Total of 17% of respondents self-reported using telehealth because of the pandemic. Black respondents were more likely than Whites to report using telehealth because of the pandemic (OR, 1.42; 95% CI, 1.07–1.88).</td>
</tr>
<tr>
<td>Campos-Castillo C &amp; Laestadius (2020) (57)</td>
<td>Cohort study</td>
<td>USA</td>
<td>10,541 USA-wide survey respondents</td>
<td>What are the characteristics of patients posting COVID-19 related messages on social media</td>
<td>Logistic regression</td>
<td>Respondents who identified as black (OR 1.29, 95% CI 1.02-1.64; P = .03), or Latino (OR 1.66, 95% CI 1.36-2.04; PP = .03) had higher odds than respondents who identified as white of reporting that they posted COVID-19 content on social media.</td>
</tr>
<tr>
<td>Chunara R, et al (2020) (50)</td>
<td>Cohort study</td>
<td>USA</td>
<td>140,184 patients</td>
<td>What are the characteristics of patients that use telemedicine</td>
<td>Descriptive statistics and logistic regression</td>
<td>Black patients nearly half as likely as white patients to access care through telemedicine (OR 0.6 times (95% CI: 0.58–0.63)</td>
</tr>
<tr>
<td>Eberly, et al. (2020) (53)</td>
<td>Cohort study</td>
<td>USA</td>
<td>2,940 cardiovascular outpatients</td>
<td>What are the characteristics of patients that complete a telemedicine consultation</td>
<td>Logistic regression</td>
<td>Being female and being non-English speaking were independently associated with less telemedicine use.</td>
</tr>
<tr>
<td>Gabbiadini et al 2020 (56)</td>
<td>Cross-sectional</td>
<td>Italy</td>
<td>465 respondents</td>
<td>Whether the use ICT promoted perceptions of social support (mitigating the psychological effects of lockdown)</td>
<td>Separate multiple and simple regression models</td>
<td>The amount of technology use was a significant predictor of perceived social support. (OR 2.40, p &lt; 0.02, 99% CI −0.01, 0.31).</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Country</td>
<td>Sample Size</td>
<td>Population Characteristics</td>
<td>Research Question</td>
<td>Methodology</td>
</tr>
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<tr>
<td>Hughes et al (2020) (49)</td>
<td>Mixed methods</td>
<td>UK</td>
<td>156 high-risk individuals and a further 1,217 vulnerable patients over the age of 70</td>
<td>Can medical students (general practitioner trainees) use tele-consultations to assess the needs of patients and support digital access to healthcare</td>
<td>Descriptive statistical analysis. Thematic analysis of conversations issues arising, no theoretical framework named</td>
<td>A total of 22% high risk patients and 44% of vulnerable patients reported connectivity issues. Participants reported that they were confident in ordering medication online.</td>
</tr>
<tr>
<td>Runfola et al (2020) (55)</td>
<td>Cohort study</td>
<td>Italy</td>
<td>33 Bariatric outpatients</td>
<td>What are the characteristics of patients that completed a telemedicine consultation</td>
<td>Categorical data was compared using the chi-square test. Continuous variables compared using the Student's t test.</td>
<td>57.6% (19) completed the consultation. No significant differences were found between participants and non-participants in terms of age and gender.</td>
</tr>
<tr>
<td>Weber et al (2020) (51)</td>
<td>Cohort study</td>
<td>USA</td>
<td>52,585 patients diagnosed with COVID-19</td>
<td>What are the characteristics of patients that access care by telemedicine, ER or office visit</td>
<td>Descriptive statistics and multinomial regression analysis</td>
<td>White patients had the highest predicted probabilities of using telehealth (46.7%) Black and Hispanic patients over 65 have lowest predicted probability (11.3%)</td>
</tr>
<tr>
<td>Whaley et al (2020) (52)</td>
<td>Cross-sectional</td>
<td>USA</td>
<td>Data from 5.6 to 6.8 million US individuals with employer health insurance between 2018, and 2020,</td>
<td>What are the characteristics of patients that use telemedicine.</td>
<td>Logistic regression</td>
<td>Patients living in post codes with lower-income or majority racial/ethnic minority populations had lower rates of adoption of telemedicine; ≥80% racial/ethnic minority post code: -71.6 per 10,000 (95% CI, -87.6 to -55.5); 79%-21% racial/ethnic minority post code: -15.1 per 10,000 (95% CI, -19.8 to -10.4).</td>
</tr>
</tbody>
</table>
Digital Access

We identified one study concerned with the access of digital technologies, specifically the reliability of internet connectivity. It was conducted in United Kingdom (UK) primary care as part of a study whose overall aim was to explore whether vulnerable patients might be usefully supported by tele-coaching in the use of digital health technologies, in this instance by General Practitioner (GP) trainees (49). As part of these conversations a direct question was asked around internet connectivity and the authors reported that 22% of high-risk patients and 44% of vulnerable patients reported issues (49).

Digital Literacy

A total of seven studies addressed the domain of Digital Literacy and in particular an individual’s digital skill set, specifically in relation to the ways in which they accessed care. All provided comparisons of use between groups using descriptions of demographic characteristics that included age, gender and ethnicity (58). Two studies used routinely collected electronic health data, though were conducted independently of each other at two different sites in New York (United States) (50, 51). The first used data gathered from patients at New York University Hospital collected over a 6-week period to determine whether they had received their COVID-19 diagnosis at an office visit or via video consultation. The authors described that the digital infrastructure of the service was well resourced and established and therefore attributed the reduced utilization of telemedicine by black patients to factors unrelated to the digital capacity of the facility (50). The second study set in New York was also situated within a large health care centre and again compared the means of accessing healthcare between ethnic groups within the early months of the pandemic (51). They found that black and Hispanic patients were more likely to visit the Emergency Room or arrange an office visit than use telehealth than their white or Asian counterparts (51).
this instance the authors recognised that the more extensive use of ER may be due to the disproportionate number of ethnic minorities that experienced severe COVID-related symptoms (51). Another study set in the United States compared the use of telemedicine amongst commercially insured patients from 2018 through to 2020 (52). In doing so they explored differences in both the nature of the care they received and the means of access in the first two months of the pandemic and described that though there was an increase in telemedicine it did not make up the shortfall in the number of visits in comparison to the usual levels of assessing preventative or elective care amongst ethnic minorities (52). Campos-Castilo and Anthony conducted a secondary analysis of a national survey a secondary analysis of cross-sectional survey data from the Pew Research Center’s American Trends Panel, which is a national, probability-based online panel of adults (18 or older) living in US households exploring self-reported use of telemedicine and following adjustment for socio-economic status (SES), age and perceived level of threat to their health from the pandemic (no threat, minor, or major) (54). They found black patients were actually more likely to contact care providers using ICTs if they perceived their health was threatened by the virus (54).

Two studies specifically explored whether there were differences in the characteristics of patients fulfilling pre-arranged or routine video-consultations during the pandemic. One of these studies was also set in the USA and compared the characteristics of cardiovascular patients that ‘attended’ teleconsultations and found no differences in cancellation rates based on race, ethnicity, or household income. However, differences between genders were observed with those completing telemedicine tending to be male and older (53). In Italy Runfola et al explored the utilisation and subsequent satisfaction with video consultations amongst a group of bariatric patients. They found no significant differences in terms age or gender between those that succeeded or failed to complete a video call (55). However, in terms of overall numbers just under 58% of patients
fulfilled the video-consultation and the authors felt that this was due to the absence of basic computer skills and a lack of self-efficacy in utilising video-call systems (55). In relation to self-efficacy the Hughes study set in the UK also assessed vulnerable patients’ confidence and ability to order medications online and reported they were comfortable and confident with the process (49).

**Digital Assimilation**

Two studies explored the use of digital technologies in relation to maintaining or interacting with a social network. One study set in Italy described how feelings of loneliness, boredom and irritability were all reduced as a result of regular utilisation of video calls, and the positive effects on maintaining meaningful relationships and mental health (56). Meanwhile in the United States another secondary analysis of the same cross-sectional survey data from the Pew Research Center’s American Trends Panel, was conducted to understand if there were differences in the characteristics of individuals that posted COVID-19 related content to social media platforms (57). The authors discovered that proportionally members of racial and ethnic minority groups and amongst these older black males, were the most likely to contribute COVID-19 related content (57).
Discussion

General findings

Our rapid review identified how pre-existing societal disparities in access to and utilisation of health-related digital technologies were accentuated by COVID-19. We identified nine studies that explored various constructs within the three domains of our digital divide framework. In relation to Digital Access, poor internet access amongst the elderly was reported (49), as regards Digital Literacy lower levels of take-up of telemedicine amongst certain communities in the United States were described particularly among black and Hispanic patients (51-54). Within the domain of Digital Assimilation one study described how face-time technology can sustain relationships amongst dislocated peer groups (55), and another how black and elderly males, previously considered a group unprepared to share health information on social media platforms were the demographic most likely to post content on the pandemic, an important consideration in understanding the emerging scepticism of the COVID vaccine in ethnic groups (56).

Strengths and weaknesses

Our search strategy was designed to capture the experiences and broader lessons that might be learnt by exploring the initial stages of the pandemic, including those of countries that had health services of sufficient maturity to initiate agile and integrated responses. We focussed on the early phase of the pandemic in order to understand the impact of the rapid changes to service delivery on those most vulnerable to the digital divide with the intention of producing timely findings that might inform service delivery in subsequent phases. That our search uncovered so few studies can be attributed to two factors relating to the pandemic; first that the research capacity of healthcare organisations would have been compromised by dealing with the exceptional demand on their services (59, 60); second that the issue of the ‘digital divide’ which had previously failed to be
considered a priority was unlikely to be addressed during the most serious public health crisis in a
generation (61).

Although our search initially uncovered numerous titles many were opinion or editorial pieces,
demonstrating how widely recognised the phenomenon of the digital divide is but also its lack of
priority as a subject for original research (24, 28, 62-64). The studies identified were conducted
within only three countries at the time of the first wave they constituted three of the top four
worst death rates from COVID-19 per capita (65, 66).

Our rapid review discovered only a small number of heterogeneous papers of limited geographic
scope which precluded data synthesis and may have introduced a degree of bias. The lack of a
theoretical underpinning in many of the papers limited generalisability (66) and that two of the
studies relied on self-reported data (49, 54) raises familiar issues regards their reliability (67).
However previous comparisons between systematic and rapid reviews have failed to find
significant differences in the outcomes they report (68, 69) and all of our included studies offered
valuable insight into how the digital divide was magnified by the changes to health delivery in the
early stages of the pandemic.

Specific findings

Digital Access

The Hughes paper provides the latest example of how discrepancies in reliable internet
connectivity continue in England (49, 70) findings which were corroborated by the most recent
surveys of digital access conducted in the UK which found that nearly 7% of homes in England and
Wales did not have a reliable internet connection (71, 72), a lack of connectivity that disproportionately affected the elderly, those of lower socio-economic status, and the disabled (71-74).

Despite the calls to harness digital technologies on a global scale (5, 75-77), these also need to address the stubborn differences in digital access that remain within the developed world where significant divisions in digital connectivity and utility remain and continue to affect the most vulnerable members of society (8, 75, 78-86). The pandemic prompted broader acknowledgement of these differences in several health economies where a number of initiatives were introduced (64, 87). For example in the UK broadband providers lowered prices and reduced data caps for the vulnerable (88), and in the United States roving buses were used to provide Wi-Fi access for unconnected communities (89).

**Digital Literacy**

The patterns in digital literacy relating to socio-economic status, age, or race described in four of the studies we identified (50, 51, 53, 54) have been observed for nearly three decades (8, 18-20, 73, 74, 90-92). However, prior to the pandemic, using traditional methods of in-person access did not hold the same degree of risk as during a pandemic where airborne transmission led to widespread recommendations to minimise social contact(36). This may be due in part to variations in individual perception of risk influenced by personal experience, social values, and the attitudes of friends and family (93). It also reflects the resistance of the digital divide to intervention. A number of previous attempts have been made to connect less technologically enabled patients to the appropriate care (63, 94, 95). However, the non- adoption and abandonment of telehealth technologies by the intended users is common (96-99) complicated by the influences of the
provider organisation and the design and compatibility of the intervention (100). Self-efficacy, patient activation and motivation are also critical yet underexplored components of the uptake of digital technology (101) as are the impact of patients’ knowledge of their condition; the expectations of the care they should receive, their social situation and the resources at their disposal (102).

In attempting to unpick this complexity a number of theoretical frameworks have been developed intended to support adoption and produce transferable learning for a range of digital innovations (103). There have also been calls for greater patient and public involvement in designing and developing digital healthcare to ensure the needs and preferences of the full range of patients are incorporated (104, 105).

Digital Assimilation

For over a decade the internet has been recognised as a key source of health information for the public and patients, yet the precise role of social media in the communication of health-related information is less clear (106). Although limited, evidence tended to suggest that sharing health information online was favoured by the young (107) and was less so amongst the elderly or those of lower socio-economic status (SES) (108). However, one study we found described how older black males were more likely to share information about COVID-19 through social media channels than other demographic groups (57). This may in part be due to the growing reluctance amongst black and ethnic minority groups to trust information provided by healthcare professionals or the mainstream media (109-111). That highlights how the growing consumption of health information through a largely unregulated network of social media platforms can have serious repercussions for
public health (111-115). This is of particular concern when placed in the context of the growing scepticism about the COVID-19 vaccine in minority communities (116, 117).

Despite the potential for spreading misinformation the work by Runfola observed benefits for mental health from the use of face-to-face digital contact during the pandemic (55) and related work found benefits from the introduction of an online blog tailored for psychiatric patients (118). The last five years has seen a growing realisation that the responsible use of social media can be an effective means of alleviating depression and social isolation and improve mental well-being (119-121). In particular the utilisation of face-time technologies has been shown to increase and enhance social interactions (122) and engagement (123). During the pandemic these benefits were recognised by the UK government in their scheme that provided free tablets to care homes to help connect isolating residents with their families and loved ones (124).

Conclusions

The rapid incorporation of digital technologies into mainstream health care delivery due to the COVID pandemic was widely understood and accepted by patients in the developed world unwilling to breach social distancing advice. However, not all patient groups were either willing or able to utilise the digital services made available nor maximise the reported benefits of face-time technology to alleviate the effects of isolation. Our findings provide further evidence that patient engagement with any model of digital healthcare is vulnerable to complex socio-political factors. If more are to reap the potential benefits of digital healthcare then not only are acknowledged improvements in infrastructure needed but also concerted efforts to train, equip and motivate all patients in its use.
Ethics statement

Ethical approval was not required

Data Sharing Statement

No additional data is available for sharing.

Funding

The work was not externally funded

Conflicts of interest

There are no conflicts of interest

Author’s contribution

IL was responsible for the conception of the work and the design of the review. IL and SG reviewed the articles and IL led the drafting of the article with input from SG and DS. SG and DS all provided critical revisions. The final version was drafted by IL and approved by SG ad DS.

Statement

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4. Rimmer A. Technology will improve doctors’ relationships with patients, says Topol review. BMJ. 2019;364:l661.
8. van Deursen AJAM, van Dijk JAGM. The first-level digital divide shifts from inequalities in physical access to inequalities in material access. New Media & Society. 2018;21(3):354-75.


71. Sweney M. Slow digital services are marginalising rural areas, MPs warn. The Guardian. 2019.


Figure 1 PRISMA diagram

81x60mm (300 x 300 DPI)
Search terms

#1 ‘COVID 19’ or ‘pandemic’ or ‘COVID’

#2 ‘digital health’ or ‘telemedicine’ or ‘remote access’

#3 ‘digital divide’

#4 #1 and #2 and #3
<table>
<thead>
<tr>
<th>Section and Topic</th>
<th>Item #</th>
<th>Checklist item</th>
<th>Location where item is reported</th>
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<tbody>
<tr>
<td>TITLE</td>
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<td>Title</td>
<td>Identify the report as a systematic review.</td>
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<tr>
<td>ABSTRACT</td>
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<td>See the PRISMA 2020 for Abstracts checklist.</td>
</tr>
<tr>
<td>INTRODUCTION</td>
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<td>Describe the rationale for the review in the context of existing knowledge.</td>
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<tr>
<td>METHODS</td>
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<td>Provide an explicit statement of the objective(s) or question(s) the review addresses.</td>
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<tr>
<td>Eligibility criteria</td>
<td>5</td>
<td>Specify the inclusion and exclusion criteria for the review and how studies were grouped for the syntheses.</td>
<td>Page 6</td>
</tr>
<tr>
<td>Information sources</td>
<td>6</td>
<td>Specify all databases, registers, websites, organisations, reference lists and other sources searched or consulted to identify studies. Specify the date when each source was last searched or consulted.</td>
<td>Page 6</td>
</tr>
<tr>
<td>Search strategy</td>
<td>7</td>
<td>Present the full search strategies for all databases, registers and websites, including any filters and limits used.</td>
<td>Page 6</td>
</tr>
<tr>
<td>Selection process</td>
<td>8</td>
<td>Specify the methods used to decide whether a study met the inclusion criteria of the review, including how many reviewers screened each record and each report retrieved, whether they worked independently, and if applicable, details of automation tools used in the process.</td>
<td>Page 7</td>
</tr>
<tr>
<td>Data collection process</td>
<td>9</td>
<td>Specify the methods used to collect data from reports, including how many reviewers collected data from each report, whether they worked independently, any processes for obtaining or confirming data from study investigators, and if applicable, details of automation tools used in the process.</td>
<td>Page 7</td>
</tr>
<tr>
<td>Data items</td>
<td>10a</td>
<td>List and define all outcomes for which data were sought. Specify whether all results that were compatible with each outcome domain in each study were sought (e.g. for all measures, time points, analyses), and if not, the methods used to decide which results to collect.</td>
<td>N/A</td>
</tr>
<tr>
<td>Study risk of bias assessment</td>
<td>11</td>
<td>Specify the methods used to assess risk of bias in the included studies, including details of the tool(s) used, how many reviewers assessed each study and whether they worked independently, and if applicable, details of automation tools used in the process.</td>
<td>N/A</td>
</tr>
<tr>
<td>Effect measures</td>
<td>12</td>
<td>Specify for each outcome the effect measure(s) (e.g. risk ratio, mean difference) used in the synthesis or presentation of results.</td>
<td>N/A</td>
</tr>
<tr>
<td>Synthesis methods</td>
<td>13a</td>
<td>Describe the processes used to decide which studies were eligible for each synthesis (e.g. tabulating the study intervention characteristics and comparing against the planned groups for each synthesis (item #5)).</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>13b</td>
<td>Describe any methods required to prepare the data for presentation or synthesis, such as handling of missing summary statistics, or data conversions.</td>
<td>N/A</td>
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<tr>
<td></td>
<td>13c</td>
<td>Describe any methods used to tabulate or visually display results of individual studies and syntheses.</td>
<td>N/A</td>
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<td></td>
<td>13d</td>
<td>Describe any methods used to synthesize results and provide a rationale for the choice(s). If meta-analysis was performed, describe the model(s), method(s) to identify the presence and extent of statistical heterogeneity, and software package(s) used.</td>
<td>N/A</td>
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<td></td>
<td>13e</td>
<td>Describe any methods used to explore possible causes of heterogeneity among study results (e.g. subgroup analyses, meta-regression).</td>
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<td></td>
<td>13f</td>
<td>Describe any sensitivity analyses conducted to assess robustness of the synthesized results.</td>
<td>N/A</td>
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<tr>
<td>Reporting bias assessment</td>
<td>14</td>
<td>Describe any methods used to assess risk of bias due to missing results in a synthesis (arising from reporting bias).</td>
<td>N/A</td>
</tr>
<tr>
<td>Certainty assessment</td>
<td>15</td>
<td>Describe any methods used to assess certainty (or confidence) in the body of evidence for an outcome.</td>
<td>N/A</td>
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<tr>
<td>Section and Topic</td>
<td>Item #</td>
<td>Checklist item</td>
<td>Location where item is reported</td>
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<tr>
<td>Study selection</td>
<td>16a</td>
<td>Describe the results of the search and selection process, from the number of records identified in the search to the number of studies included in the review, ideally using a flow diagram.</td>
<td>Page 27</td>
</tr>
<tr>
<td></td>
<td>16b</td>
<td>Cite studies that might appear to meet the inclusion criteria, but which were excluded, and explain why they were excluded.</td>
<td>Page 9</td>
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<tr>
<td>Study characteristics</td>
<td>17</td>
<td>Cite each included study and present its characteristics.</td>
<td>Page 8</td>
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<tr>
<td>Risk of bias in studies</td>
<td>18</td>
<td>Present assessments of risk of bias for each included study.</td>
<td>N/A</td>
</tr>
<tr>
<td>Results of individual studies</td>
<td>19</td>
<td>For all outcomes, present, for each study: (a) summary statistics for each group (where appropriate) and (b) an effect estimate and its precision (e.g. confidence/credible interval), ideally using structured tables or plots.</td>
<td>Page 8</td>
</tr>
<tr>
<td>Results of syntheses</td>
<td>20a</td>
<td>For each synthesis, briefly summarise the characteristics and risk of bias among contributing studies.</td>
<td>N/A</td>
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<tr>
<td></td>
<td>20b</td>
<td>Present results of all statistical syntheses conducted. If meta-analysis was done, present for each the summary estimate and its precision (e.g. confidence/credible interval) and measures of statistical heterogeneity. If comparing groups, describe the direction of the effect.</td>
<td>N/A</td>
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<td></td>
<td>20c</td>
<td>Present results of all investigations of possible causes of heterogeneity among study results.</td>
<td>N/A</td>
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<tr>
<td></td>
<td>20d</td>
<td>Present results of all sensitivity analyses conducted to assess the robustness of the synthesized results.</td>
<td>N/A</td>
</tr>
<tr>
<td>Reporting biases</td>
<td>21</td>
<td>Present assessments of risk of bias due to missing results (arising from reporting biases) for each synthesis assessed.</td>
<td>NA</td>
</tr>
<tr>
<td>Certainty of evidence</td>
<td>22</td>
<td>Present assessments of certainty (or confidence) in the body of evidence for each outcome assessed.</td>
<td>Page 8</td>
</tr>
<tr>
<td>DISCUSSION</td>
<td>23a</td>
<td>Provide a general interpretation of the results in the context of other evidence.</td>
<td>Page 15</td>
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<td></td>
<td>23b</td>
<td>Discuss any limitations of the evidence included in the review.</td>
<td>Pages 15/16</td>
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<tr>
<td></td>
<td>23c</td>
<td>Discuss any limitations of the review processes used.</td>
<td>Page 16</td>
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<tr>
<td></td>
<td>23d</td>
<td>Discuss implications of the results for practice, policy, and future research.</td>
<td>Pages 16/19</td>
</tr>
<tr>
<td>OTHER INFORMATION</td>
<td>24a</td>
<td>Provide registration information for the review, including register name and registration number, or state that the review was not registered.</td>
<td>N/A</td>
</tr>
<tr>
<td>Registration and protocol</td>
<td>24b</td>
<td>Indicate where the review protocol can be accessed, or state that a protocol was not prepared.</td>
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<td>24c</td>
<td>Describe and explain any amendments to information provided at registration or in the protocol.</td>
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<tr>
<td>Support</td>
<td>25</td>
<td>Describe sources of financial or non-financial support for the review, and the role of the funders or sponsors in the review.</td>
<td>Page 19</td>
</tr>
<tr>
<td>Competing interests</td>
<td>26</td>
<td>Declare any competing interests of review authors.</td>
<td>Page 19</td>
</tr>
<tr>
<td>Availability of data, code and other materials</td>
<td>27</td>
<td>Report which of the following are publicly available and where they can be found: template data collection forms; data extracted from included studies; data used for all analyses; analytic code; any other materials used in the review.</td>
<td>N/A</td>
</tr>
</tbody>
</table>