What can we learn from second generation digital natives? A qualitative study of undergraduates’ views of digital health at one London university

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Abstract

Objectives: We live in a digital age and opportunities within healthcare are increasing, ranging from patient portals to wearable devices. Today’s undergraduates are second generation digital natives and are at a critical point of becoming more autonomous in their healthcare interactions. This study aims to understand their experiences of both digital and broader healthcare. This will enable a better understanding of implications for national policy, individual healthcare organisations and further research.

Methods: Undergraduates aged 18–21 participated in individual interviews or focus groups. Inductive thematic analysis was undertaken. Negative member checking and feedback on emerging themes from both participants and experts were used to increase the validity of the study.

Results: Twenty-four undergraduates participated in the study, including a high proportion of international students. Thematic analysis revealed 16 themes. Six key themes explored in this paper are: generation gap; impact on healthcare professionals (HCPs); use of technology to replace or enhance HCP interactions; use of technology to support administration/transactional activities; paper vs electronic; and personally held health and fitness data.

Conclusion: This paper highlights recommendations for the undergraduate cohort and wider populations including better articulation of benefits, making digital options more personalised and interactive, and raising awareness of dangerous ‘obsessive’ behaviour around health and fitness apps. Some of our findings challenge the assumption that this generation will automatically accept digital initiatives, including the importance this cohort continues to place on face-to-face interactions. In response, we offer some suggestions to improve awareness, utilisation and acceptance of digital health.

Keywords

Students, Internet, telemedicine, delivery of healthcare, information systems

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Introduction

We live in a digital age where we increasingly carry out everyday activities at our convenience online, such as banking and shopping. Interest in how this digital revolution can support healthcare is increasing globally.¹ If the current direction of developments in digital healthcare is to be a success, it needs acceptance and endorsement from the general ‘well’ public to become normalised in society. This area of research is rapidly changing and a new focus on developing areas of online activity has been identified and recommended.²

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Digital health is a broad term. For the purposes of this paper, we conceptualise this as the use of technology to either replace or enhance existing interactions within healthcare. This may include booking appointments online, patient portals, wearable devices, virtual appointments and other self-service platforms.

‘Digital native’ is one of several terms to describe a generation who have different expectations about life and learning, having grown up with access to digital technology. A digital native is typically someone born after 1980, while those born after 1990 are ‘secondary digital natives’ due to their exposure to Web 2.0 while growing up. It is therefore understandable to assume that undergraduates, who fit the secondary digital native profile, will be natural trailblazers and accepting of digital healthcare opportunities.

This research focused on undergraduates at a major London university. As well as being second generation digital natives, these students are also at a critical point of becoming more autonomous in their interactions with healthcare, having been removed from their existing peer groups, notably parents who remain the primary information source for healthcare. This contrasts with younger adolescents who have been more studied.

Research on younger adolescents has found that healthcare needs tend to focus on appearance; mental health; sexual health; alcohol, smoking and drugs. Studies have found that perceived barriers to primary care include embarrassment, lack of confidentiality, unsympathetic doctors, and being recognised reinforced by studies that highlight difficulties in locating confidential services. This increases with particularly sensitive issues. Should these barriers also exist for undergraduates, there may be an opportunity for digital technology to reduce them. This could encourage sensible use of health services earlier and emphasises the importance of testing the assumption that secondary digital natives will accept and utilise digital healthcare.

In England, reflecting global trends, there is an increasing focus on the use of digital systems in healthcare including the Bob Wachter review of IT implementation in the NHS, and preventative approaches are dominant in NHS policy.

While earlier digital health models were static and read only, Web 2.0 has created a range of digital interactions that are participatory. This could be quite basic and transactional (such as arranging an appointment or repeat prescription online) or more innovative (such as personally tracked data feeding predictive algorithms). Patients prefer functions that offer convenience, and if patient-facing digital services are successful, they may help to reduce pressure on healthcare systems globally to tackle prevention and long-term condition management more effectively.

While the digital divide may be reducing, as seen in survey data from NHS Choices on online health practices (Jessica Holdstock, personal communication, 2016), if those using digital services fail to see any merit, there is a risk the anticipated benefits of digital health will not be realised. There is a need to understand what the general public want, and this study will examine this area in more detail from a global undergraduate’s perspective. This is a cohort who should be comfortable with and embrace the use of digital technology for healthcare. There is an opportunity to learn from the experiences and expectations of a group we assume are happy to ‘go digital’.

Thus, this study aims to:

a. explore the experiences and expectations of both digital and overall healthcare by undergraduates to understand the context in which health services are operating;

b. understand what can be learnt from second generation digital natives that can influence the direction of national policy and future research;

c. enable individual healthcare providers to consider what specific digital initiatives they could target at undergraduates to reduce perceived barriers to healthcare.

Aims a and b provided an insight into opportunities for the wider population; however, this was not a primary intention of the research.

**Method**

**Design**

This study used focus groups and 1:1 interviews to collect qualitative data, allowing an understanding of phenomena and lived experience to be created. While focus groups consisting of friends were preferred due to the benefits of undertaking research with a group of peers, they were also undertaken with groups of strangers alongside individual interviews given recruitment time constraints. Focus groups remained small (up to 4), which is appropriate for small studies, while ensuring that the researcher could support participants and effectively transcribe verbatim. Each session explored the following:

- participants’ previous experiences with healthcare, both generally and digitally;
- ranking exercise on different mechanisms (digital and non-digital) to interact with healthcare;
- experiences of collecting, using and sharing personally held health-related data.
Sample
Participants were aged 18–21 and were studying at University College London (UCL), the largest university in London. Snowball sampling was attempted to increase study participation, and this proved successful in constructing focus groups. Twenty-six participants were recruited and invited to attend either focus groups or interviews between March and July 2016. One participant cancelled on the day of the interview, and a second participant was excluded post interview having exceeded the upper age limit by 5 years. The study aimed to recruit a minimum of 20 participants, based on the method of Fugard & Potts15.

Recruitment
Students were approached through posters displayed on student noticeboards, invitations via departmental academic administrators, and advertising in UCL’s internal student newsletter. Attempts were made to advertise via social media, but this only led to some interest from other researchers, and no participants made contact this way.
Participants who contacted the researcher were emailed a copy of the information and consent sheet and asked if they were happy to participate in a focus group or interview. Those who consented were booked into a convenient time slot. All participants received a text reminder the day before their booked slot. Participants were reimbursed £10 for their time.

Setting
The research took place on the UCL Bloomsbury campus, the main university campus. Private rooms were booked ensuring privacy and furniture was laid out to avoid items such as tables acting as a barrier. Ninety minutes were allocated per session to allow time either side of the interview for paperwork and questions.

Procedure
Participants completed a ranking exercise as part of the focus groups and 1:1 interviews. They were asked to rank their preferred communication method for booking a GP appointment, receiving an appointment reminder, a GP consultation and receiving test results. Options included the use of post, emails, SMS, telephone, portals, video and in person.
The interactive exercise increased the validity of this study, enabling participants to focus on key issues despite being ‘offline’.2 As this study asked participants to think about past experiences and what they might do in a given situation, it is difficult to know if they would actually do what they said they would do. This stimulation allowed participants to consider their options more thoughtfully.
Member checking, expert review and negative case analysis were also undertaken to increase rigour.13 The researcher ensured all members in focus groups had the opportunity to contribute to prevent results from being skewed by dominant individuals, while remaining neutral throughout to reduce bias and not lead the participants to affirm emerging themes.2,16
Full transcripts were regularly reviewed to ensure reliability.

Supporting materials
An example schedule of questions can be found in the Appendix.

Piloting
Equipment, such as recording devices and laptops, were checked. Materials were reviewed by fellow researchers and adapted accordingly. A small pilot was undertaken using both a focus group and 1:1 interview approach. Those participating in the pilot were asked to give feedback on how they found the questions and whether they had felt pressured to answer in a particular way. Pilot participants advised they had felt open to give their views without feeling pressured. Pilot participants also flagged any questions where they struggled with the meaning.
Data from the pilot study was not used for the main study. The interview schedule and scenarios evolved slightly as a result to ensure questions were being asked clearly and enabled the aims and objectives of this study to be achieved.

Ethical issues
Ethics approval (ID: 8337/001) and data protection approval (ID: Z6364106/2016/01/89) were secured from UCL. The researcher reminded participants they did not need to volunteer any medical information they did not wish to disclose, and further information on conditions was not probed. No participants became upset or distressed during interviews; however, should this have happened they would have been referred to the UCL Support Services and the session terminated.
Participants were all issued with an information and consent form prior to participation. Participants were also given the opportunity to ask questions in person before the interview or focus group began. Personal data were only available to the direct research team for reimbursement purposes and sharing emerging results. Participants were advised that short quotes
might be used to illustrate key points but that participants would remain unidentifiable. Personal data were stored on an encrypted device with the Dictaphone containing voice recordings kept securely locked away. All transcripts were anonymised. The research was compliant with the UK Data Protection Act.

**Thematic analysis**

Thematic analysis was used to analyse the data, and the researcher opted for an analytical framework approach. This systematic method and useful structure for the data supported answering research questions while also ensuring individual views were both connected across participants and maintained within their own contexts. Based on existing published guidance, the first author undertook the following steps: data familiarisation, generation of initial codes, development of early analytical framework, application of analytical framework, building of thematic map and definition of names, all data charted into framework matrix, interpretation of data and write up of scholarly report.

The development of the frame commenced during data collection, ensuring interview schedules could be iteratively adapted as necessary to test new emerging themes and to manage time constraints. The second author, an experienced researcher, validated emerging codes to support determining themes. To develop the coding frame, an inductive approach was used. Advantages include reducing bias that pre-defined frameworks may bring alongside allowing for the unexpected. The inductive approach also allowed the specific research questions to evolve throughout the coding process.

**Public, patient, clinician and expert involvement**

Preliminary results were shared with both participants and experts for feedback. Transcripts were then revisited where discrepancies had been identified, to finalise the coding frame and update the discussion. Feedback was received from 6/24 participants and 2/12 approached experts after being given 9 working days to respond.

**Results**

**Participants**

All 24 participants were undergraduates aged 18–21. Sessions lasted between 36 and 71 minutes. The majority of participants were female (20/24) and international students who had moved to the UK for university study (13/24). Participants were from the UK, China, Germany, the Netherlands, Iran, Malaysia, Romania, Singapore, Spain and Zimbabwe. Participants studied a range of subjects, with one-third having links to healthcare (medicine, pharmacy, psychology). Table 1 presents the demographics breakdown.

| Type of session | Number | Percentage |
|-----------------|--------|------------|
| Focus group     | 15     | 63         |
| Interview       | 9      | 38         |

| Age |
|-----|
| 18  |
| 19  |
| 20  |
| 21  |

| Degree                          | Number | Percentage |
|---------------------------------|--------|------------|
| Arts and Sciences               | 1      | 4          |
| Biochemical/Chemical Engineering| 3      | 13         |
| Biochemistry                    | 1      | 4          |
| Economics and related           | 4      | 17         |
| Electronic and Electrical Engineering | 1 | 4         |
| Geography                       | 3      | 13         |
| Human Genetics                  | 1      | 4          |
| Mathematics and related         | 2      | 8          |
| Medical                         | 2      | 8          |
| Pharmacy                        | 4      | 17         |
| Psychology and Language Sciences| 2      | 8          |

| Nationality and educational background |
|----------------------------------------|
| A: UK student living in London prior to matriculating | 2 | 8 |
| B: UK student who moved to London for university study | 5 | 21 |
| C: International student who completed secondary education in UK | 4 | 17 |
| D: International student who moved to the UK for university study | 13 | 54 |

Table 1. Demographics of participants.
Participants were not asked about their general health, and none were known to be affected by any severe disease. One participant described their experience of living with a manageable long-term condition, and some described ongoing mental health issues. All other participants discussed short-term health issues such as stomach aches or colds.

While demographics are useful for overall context, to avoid making participants identifiable, we do not specify how many participants of which sort did or did not agree with something. Rather than representing a wider population, these data are designed to capture diversity around phenomena.17

Coding framework

A framework was developed consisting of 16 themes (Table 2). Themes are not entirely distinct and can overlap or interrelate with each other.

Ranking activity results

Participants completed a ranking exercise to decide what their most to least preferred options were when interacting with healthcare providers for booking appointments, receiving appointment reminders, receiving test results and having a consultation. Table 3 shows the median rankings for each category, where 1 indicates most preferred and 4 indicates least preferred option. Key results included favouring of portals for appointment booking and SMS/email for test results and reminders, but consultations were almost unanimously preferred face to face. Use of telephone calls and post were consistently the least preferred options across all categories.

Contextual themes

Some themes were cross cutting and contextual, supporting the whole framework. These are presented first while key themes are presented in the next section.

Barriers to technology in healthcare. Various barriers to technology in healthcare were perceived by the participants: this included the implications of running out of mobile data or losing signal and the associated cost or interruption to service, as well as issues around security and ease of use, particularly with regard to passwords, set up and multiple log-ins.

Choice. Closely linked to convenience (below), choice was also frequently mentioned. The type and severity/urgency of the health condition were often a big influence on how participants would want to interact with healthcare (digitally or otherwise). A desire to have options was articulated. Another key sub-theme within choice was the ability to ask questions and request further information as part of a two-way conversation, where this was warranted depending on the health condition itself.

Communication methods. Digital communication methods were preferred for more transactional activities; however face to face was preferred by an overwhelming majority when it came to a GP consultation. The participants generally did not seem to trust postal services, which were seen as slow and unreliable. While telephone calls were also not favoured (or used as a communication method in general for this cohort), they were recognised as useful for asking questions.

Convenience. All participants spoke about the importance of convenience to them when making their decisions, taking into consideration factors such as cost, time, balancing other commitments, having to queue, and hours that suited them. Participants spoke of busy lifestyles and a desire to carry out transactions whenever and wherever they wanted to.

Healthcare experiences. All participants were able to relate to previous healthcare experiences of either themselves or a family member. Most personal experiences were related to short-term isolated illnesses. This was a generally well cohort.

Healthcare expectations. While participants tended to talk about negative experiences more than positive ones, many were aware of the problems facing the NHS in England such as not enough doctors and limited flexibility. Participants generally spoke about not wanting to go to the GP unnecessarily. International students sometimes referred to the culture back home where it was not routine to go to a doctor unless you were very sick.

Expectations of HCPs. Some participants spoke about trying home remedies or searching for information because they did not feel they would be able to see a doctor quickly enough, but they still valued the opinion of a healthcare professional just in case.

Information sharing within healthcare. Some participants identified that information sharing within the NHS between individual organisations was not currently the norm, either through personal experiences or through placement work in a healthcare setting.

Motivations to use new technology. Some participants spoke about needing to know there would be
benefits to using a new technology such as it being more convenient or offering reassurance to them. Participants also spoke of peer usage and competitiveness as being two reasons why they might take up a new technology.

Key themes

Use of technology to support administration/transactional activities. Participants favoured digital technology to support booking appointments and receiving appointment reminders, especially if they could carry out the

| Theme                                                      | Codes                                                                 |
|------------------------------------------------------------|------------------------------------------------------------------------|
| Barriers to technology in healthcare                       | Effort; Faith (‘Did it work?’); Integration (e.g. with mobile device); Mobile data/cost; Security; Signal/Wi-Fi connectivity |
| Choice                                                     | Ability to ask questions; Flexibility; Options that suit you; Urgency/Severity |
| Communication mechanisms                                   | Applications; Email; Instant messenger; Message board/forum; Mobile device; Notification; Portal; Post; SMS; Social media; Telephone; Video/Skype |
| Convenience                                                | Access to NHS Care; Cost; Discreetness; Frustration; Laziness; Opening hours; Other commitments; Reliability; Response; Time/speed |
| Critical appraisal of online information                   | Aesthetics; Author(s)/source; Branding; Commercial vs NHS; Fear/danger; Information overload; Legitimacy; Multiple sources; Name of website; Opinionated/dramatic viewpoints; Self-diagnosis Risk; (Un)biased information |
| Expectations of Healthcare Professionals (HCPs)            | Access to my previous medical history; Internationally held medical history; Validation of perceived diagnosis; ‘When I go to the GP…’ |
| Generation gap                                             | Anti-telephone; Concerns re impact on older generations; Human vs Machine; In the longer term...; ‘Online as the norm’ |
| Healthcare experiences                                     | Avoidance of using healthcare; Bad healthcare experience; Following advice given; Good healthcare experience, GP registration; Health Insurance; Others’ health conditions, Own health conditions |
| Impact on HCPs                                             | Can help communications; Email overload; Help factors; Hindrance factors; Research on outcomes |
| Information sharing (held by healthcare providers)         | Consent; Control who sees data; Centralised database; certification; Integration; International students; “Officialness”; Privacy; Trust |
| Motivations to use new technology                          | Benefits to me; Competitiveness; Diagnosis; Peer usage; Reassurance |
| Healthcare providers use of tech to replace/enhance HCP interactions | Focused on “me”; Levels of awareness; May be distracted; Reliance on Wi-Fi/data; Reoccurring condition vs new problem; Physical examination; See my doctor |
| Healthcare Providers use of tech to support admin/transactional activities | Avoids having to tell receptionist; Cancellations’ Faith (Did it work?); Integration into usual media channels; Integration with own device; Levels of awareness; Other languages; Password problems; Self-identification/log-in; Simple set up; Try something new/different |
| Paper vs electronic                                        | Ability to annotate; Authored by; Confidence; Endorsed by; Environment; Official; Physically given to you by HCP; Retrieval; Storage; Transportable; Value; ‘When it crashes’ |
| Personally held health and fitness data                    | Celebrate success; Danger; Empowerment; Goal setting; Obsessiveness; Novelty/fad; Perceived usefulness; Quantified self; Sharing with friends & family; Sharing with HCP; Social media ‘self-filter’; Time consuming; Types of data/apps |
| Reasons to search for information online                   | Complemented by ‘offline’ sources (e.g. friends/family); Didn’t understand what HCP said; Difficulty accessing HCP; Empowerment to validate own thoughts; Navigate the NHS; Self-care/treatment |
activities at their own convenience on personal devices. The use of technology to support test results was a bit more complex. While digital technology was favoured, a combination of notifications and then the ability to download results from a portal seemed to be preferred. This offered the convenience of being able to retrieve results at a convenient time as soon as they were ready, rather than wasting time chasing up their results. Participants raised concerns however as to how rich the information would be and how they could ask questions. Participants were positive but stressed the importance of public facing systems needing to be slick, reliable and to contain the right level of information or facilitate reliable signposting.

Use of technology to replace/enhance HCP interactions. Participants were overwhelmingly opposed to the idea of using technology to replace their face-to-face interactions. Video conferencing was unpopular due to the feeling that they would not be appropriately assessed or that the doctor might be distracted by other things during the virtual consultation. Participants were potentially open to virtual consultations for a longer term ongoing problem, but few had experience of such a condition. Concerns were also raised with internet connectivity and mobile data allowances as a challenge. Virtual consultations were most popular when considered as an option for international students to communicate with their family doctors back home.

Paper vs electronic. Despite being second generation digital natives, this group still tended to see paper as being more official or real, while a couple of participants referred to both physical and electronic formats as being equally ‘permanent’. Concerns were often raised such as information getting lost in cyberspace, difficulty annotating, systems crashing and just generally preferring paper. One participant questioned the legitimacy of receiving something online compared with knowing it had come from their doctor. However, many of the group also indicated they were simply not used to getting information in a more digital format and therefore it may require a change in mind-set. Participants generally appeared open-minded about how they could receive their information in the future.

Generation gap. Many participants identified that they were part of a new generation who were more digitally enabled and therefore more receptive to change. Some raised concerns about how older generations would accept increasing use of technology, especially if they did not have family around to help them. This re-enforced the importance of choice, in that it was felt that older generations should be able to continue booking their appointments over the phone if they wish to do so. They did however recognise that age was not necessarily a hindrance and there were benefits to the use of technology to help with a variety of things including applications to support medication reminders. A large concern was data security and trusting of online information by older generations in their families, concerns in fact shared by many of the participants who highlighted a preference for paper. Several referred to the refusal of such family members to join online banking for that very reason. Another concern was around the use of passwords and log-ins, with several participants pointing out that they too struggled to keep up with this alongside older generations.

Personally held health and fitness data. Participants had mixed experiences of personally held health and fitness data. Many had tried various apps but had found either the novelty had worn off or it was too much effort to keep up to date with inputting data. With regard to fitness or calorie tracking, participants recognised the advantage as being able to reach a goal, either through a specific fitness task (such as running a marathon) or weight loss. The majority of participants however raised concerns about the obsessiveness and dangers of such apps, either through personal experience or behaviours of their peers. Some participants mentioned apps beyond fitness, such as tracking menstrual cycles or sleep. One participant suggested that they would find it useful to have the ability to test their own blood and check for any deficiencies.

### Table 3. Median scores.

| Activities                                      | Portal — 1 | SMS/text — 1 | Face-to-Face consultation — 1 | SMS/email — 1 |
|------------------------------------------------|------------|---------------|-----------------------------|---------------|
| Booking an appointment                         |            |               |                             |               |
| Receiving appointment reminders                | In person — 3 | Portal — 2    | Email advice — 3           | Portal — 1    |
| Consultation with HCP                          | Email/SMS — 3 | Telephone — 3 | Virtual consultation — 3   | Telephone — 3 |
| Receiving test results                         | Telephone — 3 | Post — 4      | Telephone consultation — 3 | Post — 4      |

Cowey and Potts
Participants advised they generally did not share their health or fitness data with their peers. The use of ‘social filters’ was frequently mentioned whereby if they did share information they would only share positive things. Most participants were open to sharing these data with an HCP should they be asked or if it supported some of their symptoms; however, it was unlikely they would volunteer the information to their HCP automatically.

Impact on HCPs. Some participants recognised the impact digital technology may have on HCPs. This was largely seen as beneficial; however, concerns were raised about doctors becoming overwhelmed with email traffic. The group generally felt that the impact on HCPs would be positive.

Table 4 presents illustrative quotations for each key theme.

Information searching

Two final themes emerged around information searching. Participants had varied reasons to search for information online. They seemed to have high critical awareness when it came to using online sources and treated them with caution depending on factors such as source, branding and extreme opinions.

Feedback from participants and researchers

Preliminary themes were shared with both experts and the original participants of the study for feedback to increase validity of the coding frame. Participants also confirmed they did not feel led and had been able to express their own opinions. Ambiguity over some of the terms used was flagged, for example virtual consultations could be segregated into video calls, webchats and the use of WhatsApp.

Discussion

The aims of this research will each be explored in turn.

Aim 1: Explore the experiences and expectations of both digital and overall healthcare by undergraduates to understand the context in which health services are operating

Recruiting via a university allowed a range of international views to be captured, representative of a modern, global society and with participants each bringing experiences of different cultural expectations and healthcare systems. Provision of healthcare to undergraduates must recognise and cater for a variety of expectations. Participants could easily reflect on general experiences of healthcare; however, they were not always able to speak about experiences of digital healthcare. Although they could easily articulate the use of digital technology in other areas (e.g. banking), they struggled to do the same for healthcare and indeed many participants were unaware of what was already possible (an example being the use of the Summary Care Record in the UK to identify drug allergies). They had generally not experienced the same frustrations as more frequent healthcare users, such as complications of data sharing between providers, although some expressed dissatisfaction with the GP registration process, which they considered cumbersome.

Aim 2: Understand what can be learnt from second generation digital natives that can influence the direction of national policy and future research

Many of the viewpoints expressed aligned with literature on digital natives. They are a cohort with a perceived busy and stressful lifestyle, often trying to balance lectures, part-time jobs and a busy social life. Services must be convenient, reliable and slick, otherwise they will not be adopted. Choice between options is also necessary, depending on severity of the condition, as well as opportunities to seek clarification instantly where necessary. This becomes very subjective and it is questionable whether many current healthcare models can cope with supporting such demands in their current configurations.

There were however some surprising findings in our results that challenge assumptions often made that digital natives will be automatically accepting of digital healthcare. Firstly, participants’ resistance to using virtual means to interact with healthcare professionals: the value of face to face interaction was more important than saving time and cost on travel, for example. Secondly, we note their perception of how paper documents can seem more real and important.

Finally, health and fitness tracking were not met with great enthusiasm. While the quantified self-movement is growing, including the quantification of physical activity and sleep, the success of such apps and devices remains quite limited. The attitudes of our participants mirror previous research in which tracking tended to be useful only if it was being used for action, goals or sense-making. Previous research on wearables to support healthcare have suggested younger generations are more likely to be comfortable using such devices due to their more open attitudes. However, the participants’ experiences in this study were short lived and they were seen as too much
### Table 4. Key theme quotations.

| Key theme                                      | Quotes                                                                                                                                                                                                 |
|-----------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| **Use of technology to support administration/transactional activities** | “Yeah I think I find it hard when they when I ring and they suggest a time and I’m like I can’t do that time and there’s just like no time I can’t do that either and I’m like Wednesday I can do that you know you’re just trying to get booked but you know in general when you get through and speak”  
“I think online would be best with like a virtual calendar and every hour slots because if patients were to call in sometime the time they want isn’t available and the other time the nurse suggests is not convenient for the patient then the length of the call will be very long.”  
“So test results umm I was trying to find an option where I could talk to my GP personally but there wasn’t so I guess the closest was that I could receive over the phone or an opportunity for me to ask questions back you know and get answers there and then I guess it’s more personal I guess”  
“I would really prefer if there was an online system as with phoning up first of all you don’t guarantee they are picking up there are limited hours they can pick up the phone and even if they do they probably wouldn’t give you an appointment for one or two weeks’ time so it’s quite difficult spending all the time on the phone to get an appointment for two weeks’ time” |
| **Use of technology to replace/enhance HCP interactions** | “They can’t tell that over the telephone so in a way it’s sort of jeopardising every aspect of your care kind of I know that sounds drastic (laughs) but in a way it’s signs that could be missed”  
“What if the internet doesn’t work (laughs) or what if sometimes with skype I cannot hear properly the person and they might not hear me properly so what if this kind of thing happens.”  
“Also talking to a screen it just doesn’t feel right especially when it’s something that can be like quite emotional at times and if you’re under the weather, just not having that like I don’t know not having like someone there feels a bit odd”  
“Making sure the GP has his full attention on me and not like answering the phone or dealing with some other stuff” |
| **Paper vs electronic** | “If there is portal I will use it but honestly I just prefer using my using paper. So [pause]. If there is a portal I will use that but view my results I might still print them out”  
“It’s kind of psychological because sometimes it feels like there’s a receptionist gets stuff wrong that kind of thing whereas if the doctor’s giving it to you see what they’re giving you then you know you’re getting the correct information, the relevant information, so that’s what makes it more”  
“For the first ones I like to have my results by email or by post as I like to keep a record of them so that if I need to go back again and look again to them I can do it easily”  
“Umm I personally prefer stuff on paper but if it’s something like that when you read it as a one off then electronic would be fine yeah”  
“I suppose it is a lot based on mind-set. So for me I like all my other stuff like banking and stuff I’ve gone paperless and everything because I don’t like the faff of having it in the house and needing to dispose of it and stuff but it’s not very eco-friendly and all of that stuff that goes with it. I guess over time I suppose mind-set changes” |
| **Generation gap** | “Umm I think that this is something that if something like this does go ahead then I think it won’t take long for people who are my age to transition into this but older people might be like ‘what!’ So even like my mum when she logs into something like my student finance she doesn’t understand what’s going on”  
“I think they do because the portal can be quite inconvenient for them like they don’t really know how to use it how to create a password and it can be quite troublesome if they receive an SMS because they have not received anything like that before and some of it just receive and they do not know how to reply”  
“Well there will always be people who will have problems with it I guess so for example I’m teaching my grandma how to send emails now and it’s really funny so she has these problems and doesn’t even know should I click one time or double so I guess she prefers calling or even going to the doctor” |
| **Personally held health and fitness data** | “Also it’s scary when you get obsessive with it I have a lot of friends in the same way as you [to another participant] when they’re like oh my god I’ve gained 2 kilos in the last like 6 hours”  
“‘I’m not umm unhealthy but I feel for some people it would help just to shock them into doing something’” |

(continued)
hassle. This is unsurprising when other studies have found even patients with long-term conditions may see the tracking of data as ‘work’. Overall, this cohort therefore did not feel the value outweighed the effort required of them.

It would be wrong to assume this cohort will be completely accepting of digital healthcare. That they appear to have more faith in face-to-face appointments and paper documents flies in the face of much national policy. However, we note that, in the UK context, there is considerable ongoing work promoting the sharing of relevant and useful healthcare information. It is likely this cohort have not experienced much of that as yet. Further, nearly all experiences the participants discussed were isolated episodes of ill health. The management of chronic long-term conditions is radically different.

Several studies have flagged issues with technical, logistical and regulatory challenges around new models of care for GP consultations. This includes email, virtual online consultations and text messaging. The overall consensus was that some mechanisms may work for particular patients or patient groups. For this group, activities that can be carried out through existing university portals may prove most successful, compared with new initiatives that may be perceived as risky or hard work.

Although the participants have been more dismissive of using technology than expected, they appeared open-minded and recognised mind-set changes were required, similar to how people have accepted online banking, for example. The issues raised were not unique to this age group, including concerns around cyber security and a reluctance to move away from paper.

Likewise, it could be argued they were more likely to recognise dangers around obsessiveness with data tracking. These dangers are already evident in the literature: with non-medical devices communicating assumptions on what is healthy, unhealthy obsessions with data such as ‘chronorexia’, an unhealthy obsession with ‘healthy sleeping’, and the risk of wrongly assuming a device is completely accurate.

Finally, participants indicated they would probably not share their own data with an HCP, which either indicates a lack of faith in data rigour or a fear in challenging the patient/doctor relationship. This mirrors previous studies where few patients considered their data as something useful to an HCP. Other issues include that the number of false positives may impact on healthcare professionals’ workload unnecessarily and a current lack of engagement by providers, due to a variety of reasons including a lack of usable export mechanisms. Such data tracking may therefore reinforce bad activities or behaviours without the ability to obtain a HCP’s expert opinion on the data.

Aim 3: Enable individual healthcare providers to consider what specific digital initiatives they could target at undergraduates to reduce perceived barriers to healthcare

A key barrier to healthcare for this cohort was getting a GP appointment. Participants lacked patience and would instead attend as a walk in; if they were sick, they wanted to be seen immediately and this appeared to be more important during stressful times such as examination periods. Thus, while the technology may support administrative functions, it is unlikely to support their underlying goal of seeing the GP quicker if the same appointments are offered. The opportunity to

| Key theme | Quotes |
|-----------|--------|
| "I think when you put anything on social media you kind of put a bit of a filter on yourself" |
| "I can’t be bothered to track everything you know to put in I weighed this many kilos today and I’ve eaten this much today or walked this much and I don’t even know what other kind of health aspects there are. I can’t be bothered to make it so that it’s like a big well-rounded thing but I also don’t think individually they are a good sign of health so I probably wouldn’t do that either" |
| "It became like an obsession and I noticed other people just being like what are you doing as before I eat I would have to like fill in all this information about what I was eating and it just became a bit of abnormal it was hindering everyday life I was sort of waking up and oh I did better than yesterday and I don’t know it just became sort of obsessive. I’ve got quite an obsessive personality and I was just like I need to stop" |
| Impact on HCPs | "With GP appointments so everyone gets like 10 minutes per patient if your patient is happy to have a skype thing or an email thing you can just use that 10 minutes block for that patient to have it however they want to have their appointment and it would save time because if it is an email they pretty much have one question you can just answer in one email and it’s done or like another five minutes you can use on your next patient so hopefully it would help with like waiting times" |
see a doctor quicker through virtual means is less relevant given the preference for face-to-face consultations.

Participants often spoke of visiting their GP with conditions that could be treated through self-care and over-the-counter medications. Despite showing high levels of scrutiny towards online information, they continued to feel the need for a GP to rule out anything more sinister. This presents a second barrier whereby participants struggled to make sense of the overwhelming amount of information available to them and potentially led to avoidable appointments being made. There is an opportunity to target undergraduates with more communication direct from the GP practice so they feel the information is relevant to them, such as assistance with processes like registration or health campaigns. This could include better signposting to trustworthy information sources.

Several participants raised the point that some initiatives such as video calls or portals may be more useful in managing a long-term condition; they just could not relate them to their personal experiences. There are opportunities within many healthcare organisations to explore what this means for their patients.

**Limitations**

Although this study was able to address the aims laid out, a number of limitations are recognised. Given constraints, pragmatic mitigation actions were undertaken where possible.

**Digital natives.** While this paper did not set out to prove or disprove the concept of a digital native, the researcher may have recruited those keener on digital technology than the average undergraduate. This is because the majority of participants were recruited via email or electronic newsletter and the advert asked for people who undertake other digital transactions such as online banking to get their attention. It may have excluded undergraduates less keen on using digital in general.

**Validity.** Many activities discussed were things participants had no experience of, such as booking appointments online. It is therefore difficult to know if what participants say they will do is what they would actually do. This limitation was identified at the start of the project and the ranking exercised was used as a mitigation, aimed to give participants some headspace to consider their options against one another. The use of participant and expert feedback also mitigated the impact of this.

**Ranking exercise choices.** While useful to focus participants, some of the choices that had been grouped in the activity were interpreted differently by participants.

It is also unclear if participants felt they could offer alternative options despite being invited to.

**Interviewer bias.** The researcher introducing themselves as working for the NHS may have led to participants being more positive, trying to please the researcher.

**Sample bias.** The majority of participants in this study were female (20/24) and international students (17/24); 13/17 had moved to the UK for undergraduate studies and 4/17 had completed secondary education in the UK prior to undergraduate studies. This may have led to a sample bias.

**Implications**

This was a modest study conducted at one site. Results are largely concordant with the existing literature, but a larger evidence base is needed before firm recommendations can be made, and we must exercise caution before making assumptions around digital natives. If further work replicates these findings, however, we suggest the following recommendations:

1. Transactional basic ‘self-service’ activities should continue to be developed, with the benefits better articulated to patients so they are aware of and can take advantage of their options.
2. Patient-facing digital health should be compatible with smartphones and involve simple yet secure sign-up processes.
3. Health economies (both globally and locally) should be cautious not to invest in ‘solutions’ that will not be used by patients; the target patient group should be considered carefully when making such decisions.
4. Healthcare providers and universities should collaborate to consider how existing student portals can be best utilised.
5. Digitisation will work best with undergraduates when they can access additional content or resources when desired: a patient receiving test results electronically, for example, must be signposted to further information or a mechanism to ask questions and obtain answers within a sensible timeframe.
6. Campaigns and information should be delivered to this cohort in a format that is useful for them. Studies have shown the benefit of using social media to target such groups, and universities should consider more effective campaigns relating to healthcare.
7. Healthcare systems should undertake further work with second generation digital natives to help break down barriers in digital healthcare. This group can provide a valuable bridge across generations given their greater experience of digital services generally.
8. Considering healthcare beyond undergraduates, our findings support the view that increased regulation is required to protect both patients and HCPs when using more innovative methods to communicate. In addition, healthcare providers need to carefully consider to whom they offer such services. Patients not wanting to communicate electronically with their HCPs have been evident in other studies, and we must ensure, as new initiatives are rolled out, they are what patients want.

9. Further research is recommended given many participants highlighted a trend around obsessiveness with personal health and data tracking, to understand whether such applications are increasing eating disorders or other anxieties around ‘the quantified self’. This is evident throughout undergraduates and their peer groups. Universities should be aware of this and be prepared to educate their students of the potential dangers.

**Conclusion**

This paper highlights an opportunity for healthcare systems globally to reach out more effectively to undergraduates. More collaborative work needs to be done to understand why they are not convinced of the benefits and to break down barriers to use. This study highlights how implementing technology alone is not the default answer if the advantages are not obvious, even to our digital natives. If we fail to do this, we risk switching on more digital solutions that fail to deliver benefits to patients, clinicians or healthcare systems.

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Appendix: Example schedule of questions

- How many of you have moved away from home/your parents when you started at UCL?
- How many of you have registered with your University GP?
  - Prompt: How easy/difficult was it?
  - Prompt: How did you feel?
- Please think about your last experience at the GP. Did you search for information before approaching your GP?
- How did you book your appointment with your GP?
- Did you have to give information to a receptionist/administrator first?
- Did you receive any correspondence before or after your appointment?
- Do you think the NHS uses technology well?
- Do you use it [technology] in other aspects of your life, e.g. online banking?
- Ask participant to complete ranking exercise
  - Prompt: How did you decide the top 3/bottom 3?
  - Prompt: Was anything missing what you would have liked to have seen?
- Did you share stuff about your health or fitness on social media?
  - Prompt: What do you think of people that do?
- Do any of you currently track your health or fitness data?
  - Prompt (if applicable): Did you do anything differently?
- Would you share these data with your GP?
- Do you think digital technology will affect the way people think about their health and wellbeing?
- Is there anything else you would like to discuss further that we haven’t yet covered?