Original Article

Variations in Demand across England for the Magnetic Resonance-Linac Technology, Simulated Utilising Local-level Demographic and Cancer Data in the Malthus Project

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Abstract

Aims: Cancer incidence varies across England, which affects the local-level demand for treatments. The magnetic resonance-linac (MR-linac) is a new radiotherapy technology that combines imaging and treatment. Here we model the demand and demand variations for the MR-linac across England.

Materials and methods: Initial clinical indications were provided by the MR-linac consortium and introduced into the Malthus radiotherapy clinical decision trees. The Malthus model contains Clinical Commissioning Group (CCG) population, cancer incidence and stage presentation data (for lung and prostate) and simulated the demand for the MR-linac for all CCGs and Radiotherapy Operational Delivery Networks (RODN) across England.

Results: Based on the initial target clinical indications, the MR-linac could service 16% of England’s fraction burden. The simulated fractions/million population demand/annum varies between 3000 and 10 600 fractions/million at the CCG level. Focussing only on the cancer population, the simulated fractions/1000 cancer cases demand/annum ranges from 1028 to 1195 fractions/1000 cases. If a national average for fractions/million demand was then used, at the RODN level, the variation from actual annual demand ranges from an overestimation of 8400 fractions to an underestimation of 5800 fractions. When using the national average fractions/1000 cases, the RODN demand varies from an overestimation of 3200 fractions to an underestimation of 3000 fractions.

Conclusions: Planning cancer services is complex due to regional variations in cancer burden. The variations in simulated demand of the MR-linac highlight the requirement to use local-level data when planning to introduce a new technology.

Key words: Demand modelling; health services needs and demand; health services research; MR-linac; radiotherapy; Radiotherapy Operational Delivery Network

Introduction

Cancer is a leading cause of mortality, in both the UK and worldwide, with 367 167 new diagnoses in the UK every year (2015-2017) [1]. Incidence rates have increased by 7% in the past decade [1] and the number of cases is predicted to rise by a further 1.6%/year until 2035 [2]. Cancer survival is also increasing, and has doubled in the past 40 years, with an estimated 50% of patients diagnosed with cancer in the UK now surviving for over 10 years [3]. This is largely due to successful advances in our ability to diagnose and treat the disease [4]. It is estimated that about 40% of patients who are cured of their disease have a treatment that includes radiotherapy [5]. However, reports in 2007 [6], 2012 [7] and 2014 [8] indicated that provision of radiotherapy in the UK falls short of demand [6,9]. Initiatives and models have been introduced in an attempt to quantify the gap between current levels of conventional radiotherapy provision and estimated demand at both a national [10,11] and local level. One mathematical model currently being used for both national and local-level radiotherapy demand simulations is the Malthus model [12]. Previous work has shown the importance of accounting for local variations in demand [13,14]. Malthus outputs showed about a three- and five-fold variation in local-level demand in terms of fractions/head of population for breast and prostate cancer, respectively [13].
As well as increasing the provision of conventional radiotherapy treatments, there has been increased utilisation of advanced radiotherapy modalities, including image-guided radiotherapy, intensity-modulated radiotherapy and proton beam therapy [15,16]. A common goal is increasing the number of adaptive radiotherapy treatments, defined as ‘aiming to customise each patient’s treatment plan to patient-specific variation by evaluating and characterizing the systematic and random variations through image feedback and including them in adaptive planning’ [17]. One attempt to achieve this is the development of simultaneous magnetic resonance imaging (MRI) and radiotherapy treatment delivery [18]. Examples include the MRIdian [19] (ViewRay, Cleveland, Ohio, USA), the Australian MR-linac system [20], the Linac-MR [21] (University of Alberta, Canada) and the Unity MR-linac system developed by Elekta (Stockholm, Sweden) and Philips (Amsterdam, the Netherlands) in partnership with The University Medical Centre, Utrecht [22]. At the time of writing, in England there are two Unity machines in operation, one at The Christie NHS Foundation Trust and one at The Royal Marsden NHS Foundation Trust, and one MRIdian at GenesisCare Oxford.

Technically, the MR-linac could be used to treat any current clinical radiotherapy indication. However, the MR-linac is more expensive than a conventional linac (£5 million [23] versus £1.6 million [24]). Therefore, the clinical target must be carefully selected for maximum impact. In times of budget limitations and demand for return on investment, it is important that any new technology or treatments should be introduced in a cost-effective manner [25–27]. This may be achieved by ensuring that facilities and resources are distributed to maximise their utility. It is important to identify specific regions within a country where there is the demand for the technology and the infrastructure is in place to deliver it. Predicting local demand for nascent services and prioritising high-quality local data above nationally averaged data is one way to accomplish this aim.

Here we aim to determine and show the variation in regional demand for the MR-linac system to aid the implementation of the technology and maximise both its utility and cost-effectiveness. This will be achieved by encoding consensus clinical indications, provided by the MR-linac consortium [28], into the Malthus model [12].

Materials and Methods

This study adapted the existing evidence-based clinical decision trees (CDTs) in the Malthus model [13] with the addition of some consensus-derived indications for the MR-linac. Although papers, such as Corradi et al. [29], have presented the potential initial applicability of a MR-guided radiotherapy solution across a broad range of different clinical sites, the clinical indications used for this study were determined by the MR-linac consortium in conjunction with international experts [30] and are shown in Table 1. The MR-linac consortium was formed in 2012 and consisted of seven international institutes installing clinical prototypes of Elekta’s MR-linac and technical partners. The member base included radiation oncologists, physicists, technologists, engineers, dosimetrists, radiation therapists, researchers, epidemiologists, radiographers and statisticians. The clinical indications were selected through the collaboration of the whole consortium and were based on expected clinical benefits, such as increased local control, decreased toxicity and a better quality of life [30].

The existing Malthus CDTs contain around 2000 evidence-based clinical decisions relating to conventional radiotherapy [32]. The introduction of MR-linac treatment indications impacted on six CDTs — prostate, central nervous system, head and neck, non-small cell lung cancer, oesophagus and pancreas (see Table 1).

Most of the MR-linac indications align with current stage-based radiotherapy indications included in Malthus, such that an entire stage grouping or treatment indication could be remapped from conventional radiotherapy to an MR-linac indication. Where divergence occurred from the current CDTs, new data were sourced, as indicated below, to ensure the appropriate proportions of patients were allocated to the new MR-linac indications. Specifically, Malthus separates glioma into low grade (I and II) and high grade (III and IV), whereas the MR-linac target indications are grade II, III and IV gliomas. Here we sourced data from published literature [31], with input from clinicians with expertise in this area to estimate the proportion of low-grade gliomas that are grade II. For the main simulations, we assumed that the MR-linac indications for all cancer sites would be treated with the same numbers of fractions as conventional radiotherapy.

Given the utility of the MR-linac for the implementation of hypofractionation, but current lack of clinical evidence, we will use two hypothetical examples of hypofractionation to show the effect it may have on the demand for the MR-linac. We will simulate the effects of 15-fraction hypofractionation in stage III lung cancer [33] and three-fraction extreme-hypofractionation in intermediate-risk prostate cancer [34]. We have assumed that if a patient is eligible to receive radical radiotherapy they are also eligible to receive hypofractionated radiotherapy. Therefore, for the hypofractionation trial we will simply change the conventional number of fractions to the hypofractionated number.

The Malthus model has been previously described [12,35]. In brief, Malthus routes a population of virtual cancer patients, which are representative of the demographics and incidence profile in the region being simulated, through the relevant disease-specific CDTs. The CDTs are based on evidence gathered from guidelines, clinical trials, registry data, national consensus and expert opinion. The Malthus model CDT evidence base closely aligns with the Royal College of Radiologists’ dose fractionation document [36]. Malthus collects information on how many virtual patients were prescribed radiotherapy (either conventional or MR-linac) and the number of fractions prescribed during a simulation. The virtual patients traverse through the CDTs in a Monte-Carlo integration, undertaking 1 000 000 walkthroughs to ensure every clinical decision is adequately represented and the averages
from the walkthroughs taken. Data for each Clinical Commissioning Group (CCG) and cancer site have been collected to show the number of patients eligible for treatment on the MR-linac.

An initial piece of scoping work testing the suitability of using Malthus for MR-linac indications was undertaken by Sanderson et al. [37] in a single region in England for lung cancer and prostate cancer. Here, Malthus was used to create a statistically representative cohort of virtual cancer patients for every CCG within England, and also for England itself, for the year 2019. This used CCG-level incidence projections of individual cancer sites [2] and the Office for National Statistics sub-national population projections [38]. CCG-level stage presentation data were included for lung and prostate cancer [39] and 3 years of data were used (2013–2015) [40]. There were no stage data for central nervous system and head and neck tumours available at the CCG level. Likewise, the levels of incomplete data and unstaged data were also too high for oesophagus and pancreas to enable accurate forecasts at the CCG level, so the national averages were used.

Given the cost and lack of clinical experience of the MR-linac, implementation strategies should focus on the establishment of a supra-regional network of machines, such that patients travel to the nearest comprehensive centre to access treatment. In England, the new Radiotherapy Operational Delivery Networks (RODN) [41] are prime candidates for these regional networks. Table 2 shows which cancer alliance(s) contribute to each RODN. Each RODN contains at least one large tertiary centre capable of delivering a comprehensive cancer service. CCG boundaries [42] on the resulting heat maps [43] do not include the most recent boundary change between NHS Cumbria CCG and

### Table 1

| MR-linac clinical indication | Malthus cancer site | Malthus stage group | Malthus CDT branch | Adjustments to select subgroup and data source |
|-----------------------------|---------------------|---------------------|-------------------|-----------------------------------------------|
| Stage III non-small cell lung cancer | Non-small cell lung cancer | Stage 3a | Surgery – positive margin | — |
| Intermediate risk prostate cancer | Prostate | Intermediate risk | No surgery – definitive radiotherapy | — |
| T1–2, N0–2a, small volume in n2b, low risk human papilloma virus positive oropharyngeal cancer | Head and neck – oropharynx | Stage I–II | Fit for curative | Data taken from Christie database, 47% of stage III –IVB are eligible |
| Grade II, III, IV gliomas (eligible to receive standard fractionated radiotherapy [60 Gy/30 or 59.4 Gy/33] with concurrent temozolomide) | Central nervous system | High grade glioma | Radical radiotherapy | 8% of low grade reduced to 6.6% for grade II only (low = I+II) [31] |
| Locally advanced pancreatic cancer | Pancreas | Stage I | Non-resectable | Data taken from Christie database, 36% [2016] –40% [all years] of stage II eligible |
| cT2–4N0/cTxN1–3M0 oesophageal cancer | Oesophageal | Stage I | Radiotherapy | — |
| | | Stage II | All radical radiotherapy indications | — |
| | | Stage III | All radical radiotherapy indications | — |
NHS North Lancashire CCG, forming NHS Morecambe Bay CCG and NHS North Cumbria CCG. Consequently, we have assumed that the RODN boundary in the North West (between 9 and 11) also does not contain this boundary change.

Results

Overall, the simulation estimates that 4.2% of England’s cancer patients would be eligible for treatment on the MR-linac, based on the initial clinical indications provided by the consortium. This is in the context of a total predicted appropriate rate of radiotherapy utilisation (ARR), which is the percentage of simulated patients requiring radiotherapy treatment, of 40.5% (excluding retreatments) (see Table 3).

If standard fractionation schemes are applied, the MR-linac is simulated to be eligible for up to 16% of the simulated fraction burden for England. The total number of fractions simulated for the MR-linac (MR-linac fractions) is about 351 000, with lung and prostate treatments accounting for over 60% of that figure. Table 3 shows the number of fractions for the cancer sites relevant to the MR-linac and the contribution to the overall simulated fraction burden of that cancer site.

Figure 1 shows the number of standard-scheme MR-linac fractions and the number of fractions simulated for the MR-linac/million population for each CCG on a map of England. Figure 2 shows the number of MR-linac fractions and fractions/million population simulated for each RODN on a map of England.

Due to the variations in cancer stage presentation and overall differences in case-mix across England, the simulated MR-linac fractions/million virtual population/year ranges from 3000 up to 10 600 fractions/million at the CCG level. Focussing only on the cancer population, the simulated MR-linac fractions/1000 cancer cases ranges from 1028 up to 1195 fractions/1000 cases. At the RODN level, the simulated fraction demand ranges from 13 700 up to 48 200 fractions. The number of simulated MR-linac patients ranges from 520 up to 1750/network. Due to the heterogeneity of the population within England, two regions with a very similar percentage of eligible patients may not have the same MR-linac fraction demand. There are 21 CCGs with an MR-linac ARR of 4.0%, but across those 21 CCGs the fractions/million demand for the MR-linac varies from 3500 to 8600 fractions/million. These observations are consistent with previous applications of our model.

Table 2 has grouped the results into the 11 RODNs of NHS England. The table highlights the over- or under-prediction of demand if the average MR-linac fractions/million and fractions/1000 cases for England is applied to the RODNs without using any local-level data. The largest differences between a local-level simulation for a RODN and what the estimated demand if the national average fractions/million is used were an overestimation of 8400 fractions for RODN 1 and 2 and an underestimation of 5800 fractions for RODN 9. When using the national average fractions/1000 cases, RODN 1 has an overestimation of 3200 fractions and RODN 6 an underestimation of 3000 fractions.

Table 4 shows the effects of two hypothetical examples of hypofractionation that could apply to the MR-linac. If hypofractionation schemes are applied to stage III lung and intermediate-risk prostate cancer, the simulated fraction demand for the MR-linac accounts for 11% of the simulated fraction burden for England. Compared with standard fractionation schemes, the simulated demand would drop from 107 400 to 49 900 fractions for lung cancer and from 112 100 to 16 800 fractions for prostate cancer. The total number of MR-linac fractions decreases by about 45% from 351 000 to 198 200 fractions. The number of patients would remain the same.

Discussion

There are variations in cancer incidence across the country, including variations in the case-mix and stage at presentation, and these variations affect the local demand for cancer services. When planning to introduce a new technology into cancer services, the local-level demand should be taken into account. If the introduction of a new technology occurs where demand is too low, there could be insufficient patients for an adequate economic evaluation. There are research tools available, with granular cancer incidence and population data, that can model the local-level demand for radiotherapy. Although there are a few radiotherapy demand models available [10,11], here we show the use of the Malthus model to estimate the national demand and to quantify regional variations in demand for the MR-linac. This was achieved by modifying the current clinical evidence base of Malthus to include new target MR-linac indications for radiotherapy.

The target clinical indications chosen are not a definitive list of potential indications for any form of MR-guided radiotherapy. The clinical indications in this study were chosen as the MR-linac consortium has working groups focussing on those disease sites. Therefore, there is active research in those areas and specific indications could be provided to include in the modelling. The addition or removal of clinical indications could have a large influence on the potential demand, depending on both the target cancer site and the number of fractions. The removal of the prostate clinical indication in this study would reduce the number of patients by about 42%. This shows that the right balance must be struck between target clinical indication to achieve outcomes and health economic analysis to determine cost-effectiveness. By utilising a pre-existing model, such as Malthus, scenarios can be run to estimate the impact of changing clinical indications without creating a new demand model every time.

Malthus simulated the demand for the MR-linac at the national, RODN and CCG level. It estimated that 4.2% of all cancer patients could be eligible for this treatment. This translates to around 16% of the entire fraction demand across England. The fraction demand reduces to 11% with hypothetical examples of hypofractionation for stage III lung and intermediate-risk prostate cancer. Although
Table 2
Total simulated magnetic resonance-linac (MR-linac) fractions in the Radiotherapy Operational Delivery Networks (RODN) within NHS England (also showing the respective cancer alliances covered [40]), the population, the number of cancer cases and simulated MR-linac cases in each network. Also shown is the difference in a region’s simulate demand if England’s average demand figures were used in place of local-level data.

| RODN Number | Cancer alliance(s)                                      | Population | Cancer cases | Simulated MR-linac patients | Simulated MR-linac fractions | MR-linac fractions/million population | MR-linac fractions if England average used | Difference between local-level simulation and if England average is used | Fractions/1000 cancer cases | Fractions/1000 cases if England average used | Difference between local-level simulation and if England average is used |
|-------------|--------------------------------------------------------|------------|--------------|------------------------------|-----------------------------|--------------------------------------|--------------------------------------------|--------------------------------------------------------------------------------|-----------------------------|-----------------------------------------------|--------------------------------------------------------------------------------|
| 1           | North West and South West London Surrey and Sussex     | 6 772 300  | 32 245       | 1280                         | 33 815                      | 4993                                 | 22 122                                     | 8424 (−61%)                                      | 1017                        | 36 997                                       | 1174 (−22%)                                    |
| 2           | North Central and North East LondonKent and Medway     | 3 550 100  | 12 435       | 516                          | 13 698                      | 3858                                 | 23 010                                     | 3222 (−16%)                                      | 1113                        | 19 783                                       | 1195 (−3%)                                    |
| 3           | South East London Somerset, Wiltshire, Avon and Gloucestershire | 4 584 800  | 29 762       | 1173                         | 30 599                      | 6674                                 | 28 570                                     | −2029 (−7%)                                      | 1028                        | 33 121                                       | 1054 (−3%)                                    |
| 4           | Thames Valley Wessex                                   | 5 000 100  | 29 252       | 1187                         | 30 830                      | 6166                                 | 31 158                                     | 328 (−6%)                                       | 1054                        | 32 554                                       | 1724 (−12%)                                   |
| 5           | East of England                                        | 6 559 700  | 37 109       | 1751                         | 44 354                      | 6762                                 | 40 877                                     | −3477 (−8%)                                      | 1195                        | 41 297                                       | 3056 (−7%)                                    |
| 6           | East Midlands                                          | 4 214 600  | 24 228       | 1017                         | 26 893                      | 6381                                 | 26 263                                     | −630 (0%)                                       | 1110                        | 26 962                                       | 69 (−7%)                                      |
| 7           | West Midlands                                          | 5 861 500  | 33 254       | 1470                         | 38 369                      | 6546                                 | 36 526                                     | −1843 (−5%)                                      | 1154                        | 37 007                                       | −1362 (−4%)                                   |
| 8           | Lancashire and South Cumbria Greater Manchester Cheshire and Merseyside | 6 801 300  | 41 109       | 1754                         | 48 244                      | 7093                                 | 42 382                                     | −5862 (−12%)                                     | 1174                        | 45 749                                       | −2495 (−5%)                                   |
| 9           | Humber, Coast and Vale West Yorkshire, South Yorkshire, Bassetlaw, North Derbyshire and Hardwick | 5 850 400  | 34 797       | 1397                         | 38 238                      | 6536                                 | 36 457                                     | −1782 (−5%)                                      | 1099                        | 38 724                                       | 486 (−5%)                                     |
| 10          | North East and Cumbria                                  | 3 313 700  | 21 729       | 931                          | 25 389                      | 7662                                 | 20 649                                     | −4740 (−19%)                                     | 1168                        | 24 181                                       | −1208 (−5%)                                   |
| England     |                                                        | 56 201 000 | 313 697      | 13 210                       | 350 215                     | 6231                                 | N/A                                        | N/A                               |

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hypofractionation will be a significant part of the clinical utility of the MR-linac, there is currently a lack of consensus and efficacy data on appropriate hypofractionation for each treatment indication. Therefore, although the access rate figures and simulated patient numbers are probably accurate, the fraction demand figures should be considered as an upper limit of fraction burden.

The fraction demand percentage is noticeably higher than the percentage of eligible patients because the target MR-linac indications used in this study are for radical intent only. However, the simulations will evolve and the numbers updated as evidence is generated or hypothesised for novel clinical MR-linac indications. For example, one potential MR-linac target that could increase the patient numbers to a greater extent than the fraction burden is oligometastatic disease.

The estimated demand for the MR-linac does vary across England, due to the regional variations in overall cancer burden, cancer case mix and stage at presentation. For the MR-linac, the initial lung and prostate clinical indications targeted only a limited stage/risk and therefore the local-level stage at presentation has a strong effect on demand. The overall patient demand ranges from 3% of CCG cancer incidences up to 6% and the fractions/million demand ranges from 3000 up to 10 600. When analysing simulated fraction demand as a proportion of cancer cases, the difference in demand is less marked, ranging from 1020 to 1170 fractions/1000 cases.

Without correction for regional cancer burden, demand overestimates of up to 60% would be observed in the south of England, especially London with its younger age structure. Underestimations of up to 20% would be observed in

Table 3
Simulation results for standard fractionation schemes showing the magnetic resonance-linac (MR-linac) targeted six cancer sites, and the figure for all cancers for comparison with the total radiotherapy demand, extracted from a simulation of England (2019) covering all of the Malthus 23 cancer groups

| Cancer site          | Appropriate rate of radiotherapy | % patients receiving conventional radiotherapy | % patients receiving MR-linac | Average fractions/MR-linac patient | Total MR-linac fractions | MR-linac fractions as % of all fractions for cancer site |
|----------------------|---------------------------------|-----------------------------------------------|------------------------------|-----------------------------------|--------------------------|--------------------------------------------------------|
| Central nervous system | 68%                             | 34%                                           | 34%                          | 29.6                              | 46 000                   | 65%                                                   |
| Head and neck         | 81%                             | 68%                                           | 13%                          | 34.2                              | 46 300                   | 17%                                                   |
| Lung                  | 61%                             | 52%                                           | 9%                           | 32.3                              | 107 400                  | 36%                                                   |
| Oesophagus            | 30%                             | 15%                                           | 15%                          | 27.3                              | 31 600                   | 84%                                                   |
| Pancreas              | 15%                             | 12%                                           | 3%                           | 28                                | 7600                     | 34%                                                   |
| Prostate              | 51%                             | 37%                                           | 13%                          | 20                                | 112 100                  | 29%                                                   |
| All cancer sites      | 40.5%                           | 36.3%                                         | 4.2%                         | –                                 | 351 000                  | 16%                                                   |

Fig 1. Number of fractions and fractions/million population simulated for each Clinical Commissioning Group in England for the magnetic resonance-linac (MR-linac), based on the clinical indications provided by the MR-linac consortium.
northern England, with a lower population density and a larger proportion of elderly patients. The variations are reduced by focussing on the proportion of cancer cases belonging to an MR-linac indication group and the stage presentation by using fractions/1000 cancer cases. There is still a ±10% over-/underestimation if a national average is used. The patterns are similar to those observed when using fractions/million, with overestimations typically seen in the south of England and underestimations seen in northern England.

This level of variation shows the need for local-level planning of services utilising local-level data, especially when looking to install a new technology that only targets specific tumour sites. Even when CCGs are grouped into RODNs and an averaging effect might be expected, the variation still occurs. There is still a maximum two-fold difference in the demand across the RODNs in England.

In general, CCG-level stage at presentation data in Malthus are reasonably complete. However, in some disease sites, the quality of available stage at presentation data is not sufficient to provide accurate CCG simulation results. In these cases, high-quality national-level stage at presentation data were used. Overall, cancer data are increasing in quality over time and data that are more granular will be included when made available. This should not affect the results as two-thirds of the total MR-linac fraction burden simulated uses the CCG-level stage presentation data.

The oropharyngeal cancer CDT uses the most assumptions. Local (but comprehensive) data had to be used to determine the proportions of the Malthus stage groupings (I–II, III–IVB and IVC) that were MR-linac eligible, as the target indications were TNM based. Oropharyngeal cancer contributed around 13% to the total MR-linac fraction burden and it has the highest number of fractions/MR-linac patient. It would be recommended that a hospital use its

| Cancer site     | Standard fractionation regimen | Hypofractionation regimen |
|-----------------|--------------------------------|---------------------------|
|                 | Average fractions/ MR-linac patient  | Total MR-linac fractions | MR-linac fractions as % of all fractions for cancer site | Average fractions/ MR-linac patient  | Total MR-linac fractions | MR-linac fractions as % of all fractions for cancer site |
| Lung            | 32.3                            | 107 400                    | 36%                                                   | 15                               | 49 900                     | 21%                                                   |
| Prostate        | 20                              | 112 100                    | 29%                                                   | 3                                | 16 800                     | 6%                                                    |
| All cancer sites| —                               | 351 000                    | 16%                                                   | —                               | 198 200                    | 11%                                                   |
own oropharyngeal cancer data to determine the correct proportion of Malthus patients eligible for MR-linac treatment.

This study is designed to aid the implementation of a new technology by calculating the fraction burden and patient numbers to assist with calculations of the number of MR-linacs required in different regions. For the MR-linac, and any new technology, it is non-trivial to translate the number of fractions into throughput until the technology is fully in use with initial teething problems resolved. Any treatment activity measured now will probably require more time/fraction. In machine throughput terms this may counterbalance any operational gains made from moderate hypofractionation.

MR-linac patient-specific restrictions, such as the smaller bore size and exclusion of patients with metallic implants, were not included due to the lack of data. Therefore, the results should be treated as initial estimates rather than definitive results. There is currently uncertainty around what sites would benefit the most from hypofractionated or extreme-hypofractionated regimens and clinical research is being undertaken on this subject. The MR-linac version of Malthus is flexible enough to be adapted to new treatment indications or fractional schemes quickly, to provide updated simulations for service planning.

Conclusions

Planning the introduction of new treatment technologies across a whole healthcare system is made more complex by regional variations and services should be tailored to the local demographics to ensure adequate access to treatments. Recently, local-level cancer data have become more complete and more readily available, and should be exploited in demand simulations of healthcare services. Here, we have shown the capability of using a discrete-event simulation model, Malthus, to determine the demand for a new treatment technology at a local level. This is especially important in a resource-limited setting, where demand for treatments already outstrips the supply capabilities, during times of economic austerity with calls from governments for cost savings. The simulated variations in demand for the MR-linac should highlight the benefits of combining the local-level data with comprehensive models.

Data Availability

The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

Conflicts of Interest

T. Mee reports grants from Elekta MR-linac consortium, grants from Cancer Research UK, grants from Christie Charity, grants from UKRI, during the conduct of the study. K.J. Kirkby reports grants from CRUK ARTNET, grants from Elekta MR-linac Consortium, grants from Christie Charity, during the conduct of the study; grants and non-financial support from Varian, grants from UKRI, grants from CRUK, grants from NIHR BRC, outside the submitted work. R. Jena reports personal fees from Microsoft Research, outside the submitted work. A. Choudhury reports grants from the National Institute of Health Research, Manchester Biomedical Research Centre, grants from Cancer Research, UK, grants from Medical Research Council, UK, grants from Prostate Cancer, UK, grants from Bayer, UK, personal fees from Janssen Pharmaceutical, non-financial support from ASCO, grants and non-financial support from Elekta AB, outside the submitted work. N.F. Kirkby reports grants from CRUK ARTNET, grants from Elekta MR-linac Consortium, grants from Christie Charity, during the conduct of the study; grants from UKRI, grants from CRUK, grants from NIHR BRC, outside the submitted work.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.clon.2021.03.004.

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