

UNIVERSITY OF  
COPENHAGEN



PH.D. THESIS  
by  
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## Biological Data Science

Ancient genomics, anesthesiology, epidemiology,  
and a bit in between

*Submitted: 2022-11-22*

*This thesis has been submitted to the  
PhD School of The Faculty of Science,  
University of Copenhagen.*

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*Biological Data Science:*  
*ancient genomics, anesthesiology, epidemiology, and a bit in between,*  
2022-11-22.

*Til kvinderne i mit liv*



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# *Preface*

This Ph.D. thesis summarizes my scientific research in collaboration with the Niels Bohr Institute (NBI) and the Globe Institute, University of Copenhagen, and was funded by the Lundbeck Foundation. The research was supervised by Associate Professor Troels C. Petersen (NBI) and Assistant Professor Thorfinn S. Korneliussen (Globe Institute).

Being a cross-disciplinary project, the research presented in this thesis is multi-faceted and covers a wide range of topics with the main scope being the development and integration of a novel statistical methods and machine learning models for the analysis of large-scale biological data. The thesis is organized as follows: First I present a brief introduction to the statistical methods and machine learning models used in the thesis and then I present the research in the form of four papers, each of which reflects a different aspect of the research.

The first paper presents a novel method I developed for detecting and classifying ancient DNA damage in metagenomic samples taking the full taxonomic information into account. While the first paper focuses on the development of the statistical model in the field of ancient genomics, the second paper focuses on the use of modern machine learning models in medicine and how advanced boosted decision trees can not only improve the accuracy of identifying patients at risk of being readmitted after knee or hip surgery, but doing so in a way that is interpretable as well.

In the beginning of 2020 we all experienced how COVID-19 suddenly changed our lives and impacted our societies in dramatic ways. During this time, I worked for Statens Serum Institut, the Danish CDC, on a project to develop an agent based model capable of simulating the spread of COVID-19 in Denmark. This model is presented in the third paper and was used to inform the Danish government on how to best handle the pandemic in the early stages and the effect of contact tracing.

Finally, in the fourth paper I show how advanced Bayesian methods can be utilized to better estimate the diffusion coefficients in silencing foci in the cell nucleus with single-particle tracking experiments.



## *Acknowledgements*

First of all, I want to express my sincere gratitude to my long time supervisor, captain, and friend: Troels. You are truly an inspiration to work with. I want to thank you for opening so many doors for me, both academically, professionally, and nautically. I am looking forward to our future adventures together. I also want to thank my co-supervisor, Thorfinn. I want to thank you for introducing me to the field of bioinformatic and helping me to develop my skills in this area. I also want to thank you for your patience and guiding me through the endless amount of (near) identical biological concepts and helping me to understand the minute differences.

I have been fortunate to work with people from a wide range of backgrounds and disciplines during my Ph.D. The author lists on the papers in this thesis include a particle physicist, bioinformaticians, a clinical professor, epidemiologists, a medical doctor, a bio-physicist, a mathematician, a biologist, and the president of the Royal Danish Academy of Sciences and Letters at the time. Before anything else, I want to thank all of my co-authors for their work and contributions to these papers and for allowing me to be a part of their projects. I have learned a lot from all of you, and I hope I have been able to contribute something to your work as well.

I am thankful for all the people who have helped me with my work and listened to my complaints when I was stuck, when the code did not compile, or when the small bug was almost impossible to find (which was not a small amount of time). In particular I want to thank the people at Globe who I have spent the most time with; Rasa, Alba, and Rasmus. I also want to thank the Korneliussen Group and the people in my office for helpful advice, suggestions and discussions. This also includes Daniel Nielsen and Rasmus Ørsøe from NBI. Finally, I want to thank Mathias Heltberg for many years of fruitful collaboration and for including me in his projects.

This project would not have been possible had it not been for the Lundbeck Foundation which funded my Ph.D. In addition to the funding itself, I am grateful for the inter-disciplinary aspect of project which has allowed me to meet so many inspiring and talented people and for the freedom to pursue my own interests within the project.

I would also like to express my gratitude to Professor Guido Sanguinetti from the International School for Advanced Studies, SISSA, in Trieste, Italy, for hosting me in his group during the Winter of 2021/2022. My gratitude also goes out to

Kosio, Sara, Max, Romina, Noor, Viplove, Anne-Marie, and all the other wonderful people I met during in Trieste. Thanks for making my stay in Italy so enjoyable and for welcoming me in a way only non-Danes can do.

I want to thank my friends for always being there for me. A special thanks to my friends from NBI and Borchens who I know I can always count on, whether or not that includes a trip in the party bus of the Sea, taking Artemis out for a sail, or board games and beer. Thank you for always being there. I also want to thank my family, especially my parents for their unconditional support and encouragement. I am grateful for the opportunities they have given me and for the sacrifices they have made for me.

Lastly, I want to thank my future wife and mother of our child, Anna. I would not have been able to do this without you. Thank you for your patience and support. I am looking forward to our future together. I love you to the moon and back.

## *Abstract*

Basically a thesis (book?) class for Tufte lovers like myself. I am aware that `tufte-latex` already exists but I just wanted to create my own thing.



## *Dansk Resume*

Her et dansk resumé.



# *Publications*

The work presented in this thesis is based on the following publications:

- Paper 1:** **Christian Michelsen**, Mikkel W. Pedersen, Antonio Fernandez-Guerra, Lei Zhao, Troels C. Petersen, Thorfinn S. Korneliussen (2022). “metaDMG: An Ancient DNA Damage Toolkit”.
- Paper 2:** **Christian Michelsen**, Christoffer C. Jørgensen, Mathias Heltberg, Mogens H. Jensen, Alessandra Lucchetti, Pelle B. Petersen, Troels C. Petersen, Henrik Kehlet (2022). “Preoperative prediction of medical morbidity after fast-track hip and knee arthroplasty – a machine learning based approach”.
- Paper 3:** Mathias S. Heltberg, **Christian Michelsen**, Emil S. Martiny, Lasse E. Christensen, Mogens H. Jensen, Tariq Halasa and Troels C. Petersen (2022). “Spatial Heterogeneity Affects Predictions from Early-Curve Fitting of Pandemic Outbreaks: A Case Study Using Population Data from Denmark”. In: Royal Society Open Science 9.9. issn: 2054-5703. doi: 10.1098/rsos.220018.
- Paper 4:** Susmita Sridar, Mathias S. Heltberg, **Christian Michelsen** Judith M. Hattab, Angela Taddei (2022). “Microscopic single molecule dynamics suggest underlying physical properties of the silencing foci”.



# **1** *Introduction*

The primary content of my thesis is the four papers included in the thesis. This chapter is meant as a brief introduction to the background needed to understand the basics of the methods used throughout the papers. As such, this chapter is not meant to be a comprehensive guide to statistics and bioinformatics used in the papers. The original research motivation supporting the funding of this Ph.D. was multi-disciplinary and the papers included in my thesis are also highly influenced by this.

In Section 1.1, I will shortly introduce the field of ancient genomics and the statistical methods used to identify ancient DNA will be explained. Paper I, see Chapter 2, utilize modern Bayesian methods to classify which species are ancient, and which ones are not. Bayesian methods are great when possible, however, they also rely on some statistical model being defined. In the case of Paper I, the model is a Beta-Binomial distribution combined with an exponential-decay damage model.

Sometimes the model is not known and the data generating process has to be inferred by other means. This is the case in Paper II, see Chapter 3, where we utilize machine learning methods to extract this information. This paper deals with estimating the individual risk scores for each patient being re-hospitalized after a knee or hip operation. Section 1.2 introduces the reader to basic classification with machine learning models.

While the former two papers are based on real life data, Paper III, see Chapter 4, concerns the development of a new agent based model for COVID-19. The model is based on the SIR model, but with a more detailed description of the disease and the transmission process. The model is used to simulate the spread of the virus in Denmark and to estimate the effect of contact tracing. The model is also used to simulate and predict the spread of the “alpha” variant of COVID-19 in Denmark. Section 1.3 introduces the reader to the basics of agent based models.

Finally, the method of Bayesian model comparison of different diffusion models is introduced in Paper IV, see Chapter 5. In particular, this paper deals with different mixture-models of independent Rayleigh-distributions, and how they can be used to extract important information about the underlying diffusion processes of a polymer bridging model in cell nuclei, see Section 1.4.

### 1.1 *Ancient DNA and Bayesian Statistics*

Until the mid 1980s, studies within archaeogenetics was limited to analysis of fossilized samples of plants, animals or other species (Parducci and Petit, 2004). Following the first successful recovery of ancient DNA from 5000 years old ancient Mummies, it was shown that it was indeed possible to extract and sequence DNA (Pääbo, 1985a; Pääbo, 1985b) This discovery, along with a dozen other pushing the boundary for what is scientifically possible with ancient DNA, led to Svante Pääbo being awarded with the Nobel Prize in Physiology or Medicine in 2022 for “his discoveries concerning the genomes of extinct hominins and human evolution” (Karolinska Institutet, 2022).

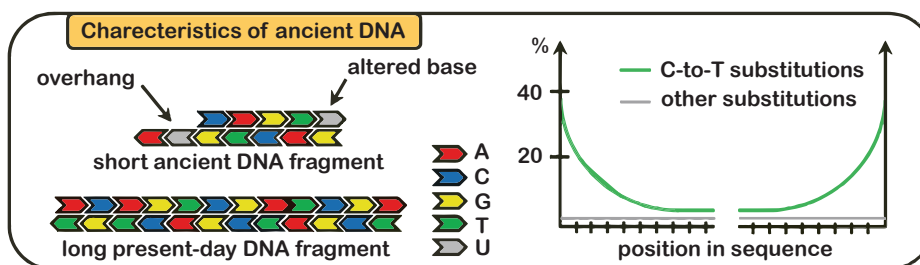
The field of ancient DNA (aDNA) was drastically changed with the invention of the Polymerase Chain Reaction, PCR, method (Mullis et al., 1986) along with the Next Generation Sequencing technology which revolutionized the speed and throughput of genomic sequencing, while decimating the cost (Slatko, Gardner, and Ausubel, 2018). This technological advance has lead to better understanding of human migration and the genealogical tree of modern humans including the previously unknown human (sub)species; the Denisova hominin (Krause et al., 2010).

Leaving the homocentric world view, aDNA also allows for the study of archaic animals. The age limitation for when aDNA can be sequenced, has in the recent years increased; in 2013 with the early Middle Pleistocene 560–780 kyr BP horse (Orlando et al., 2013) and in 2021 with the million-year-old mammoths (van der Valk et al., 2021). High-throughput sequencing not only allows for the sequencing of single genomes – like single humans, animals, or plants – but also for sequencing of entire communities of organisms, so-called metagenomics. By analysing environmental DNA (eDNA) from a set of samples, one can survey the rich plant and animal assemblages of a given area and at a specific time in the past. Our new paper published in Nature shows it is now possible to perform metagenomic sequencing on environmental DNA that is 2 million years old, see Appendix A. This is a direct application of the statistical method developed in Paper I, see Chapter 2, showing that metaDMG can help to push the boundary of what is possible with ancient DNA.

Ancient DNA is difficult to work with since it often contains only a limited amount of biological material due to bad preservation, leading to low endogenous content with high duplication rates, making high-depth sequencing difficult<sup>1</sup> (Renaud et al., 2019). In addition to this, the DNA is often highly degraded. In particular, the two prominent issues with aDNA is fragmentation and deamination (Dabney, Meyer, and Pääbo, 2013; Peyrégne and Prüfer, 2020). Fragmentation

<sup>1</sup> Genotype likelihoods are often used to alleviate the problem of low-coverage data (Nielsen et al., 2011)

refers to the fact that the DNA is broken into very short fragments, often with a size of less than 50 bp. A consequence of this, upon alignment, is low mapping quality, multimapping, and reference bias, which can somewhat be mitigated by the use of variant graphs (Martiniano et al., 2020). Deamination is a process in which cytosine (C) in the single-stranded overhangs in the end of the DNA molecules is often hydrolyzed to uracil (U) which is read as thymine (T) by the DNA polymerase. This particular type of postmortem damage is known as cytosine deamination, or C-to-T transitions, and is one of the main reasons behind nucleotide misincorporations in ancient DNA (Briggs et al., 2007). Due to the short fragment sizes in ancient DNA, the fragments will often contain overhangs with over-expressed C-to-T frequency. In the case of single-genome analysis, previous solutions have been to either remove all transitions and only keep transversions, or apply trimming at the read ends (Schubert et al., 2012). For an illustration of both fragmentation and deamination of ancient DNA, see Figure 1.



**Figure 1.**  
Illustration of DNA damage. Ancient DNA is often highly fragmented with short reads compared to modern, present-day DNA, and can contain uracils (U). These uracils will then be misread as thymines (T) while sequencing leading to C-T nucleotide misincorporations. This is primarily happening at the end of the reads. Modified from (Peyrégne and Prüfer, 2020).

Currently, a handful of different methods for quantifying ancient DNA damage exists. In particular, the mapDamage 2.0 software has been the standard for how to measure ancient DNA damage in the field (Jónsson et al., 2013), however, it uses slow algorithms leading to unfeasible runtimes for large datasets. Within genetics, newer and faster methods are continuously being developed, including PyDamage (Borry et al., 2021), which tackles some of mapDamage's limitations. However within metagenomics, which studies the genetic material of all organisms, collected from an environmental sample, faster methods suited to analyse this large-scale dataset are still lacking.

In Paper I, see Chapter 2, introduces the metaDMG software which utilizes the C-to-T deamination pattern<sup>2</sup> to identify ancient DNA damage. One of the key features of this method is the beta-binomial model which allows the uncertainty of the deamination frequency to be fitted independently of the mean of the frequencies leading to improved accuracy of the damage estimation. The deamination frequencies are based on the number of C-to-T transitions,  $k$ , out of the total number of C's,  $N$ , for a given position within the fragment. The classical likelihood to

<sup>2</sup> for the forward strand and the G-to-A deamination pattern for the reverse strand

use for this type of data is a binomial distribution. The mean and variance of the binomial distribution is given by:

$$\begin{aligned}\mathbb{E}[k] &= Np \\ \mathbb{V}[k] &= Np(1-p),\end{aligned}\tag{1}$$

where  $p$  is the probability of success (a C-to-T substitution). One of the issues, however, is that the variance of the binomial distribution is proportional to the mean. The binomial distribution is thus not flexible enough to accommodate large amounts of variance in the data, so-called overdispersion (McElreath, 2020). One way to accommodate overdispersion is to instead use a beta-binomial model. The beta-binomial model is a generalization of the binomial distribution where the variance is independent of the mean. Technically, the beta-binomial model assumes that  $p$  is a random variable which follows a beta distribution  $p \sim \text{Beta}(\mu, \varphi)$  where the beta distribution is parameterized<sup>3</sup> in terms of its mean,  $\mu$ , and dispersion parameter,  $\varphi$ , (Cepeda-Cuervo and Cifuentes-Amado, 2017). The mean and variance of this beta-binomial model is then given by:

$$\begin{aligned}\mathbb{E}[k] &= N\mu \\ \mathbb{V}[k] &= N\mu(1-\mu)\frac{\varphi+N}{\varphi+1}.\end{aligned}\tag{2}$$

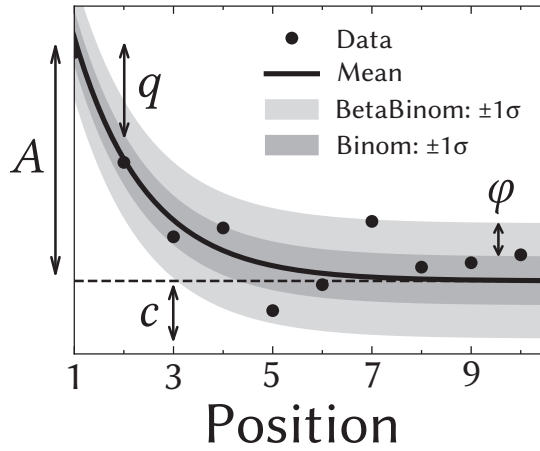
Comparing Equation 1 and Equation 2, it is seen that the variance of the beta-binomial model is no longer (strictly) proportional to the mean, but instead is a function of the dispersion parameter,  $\varphi$ , allowing for higher variance than the binomial-only model. When  $\varphi = 0$ , the variance of the beta-binomial model is  $N$  times larger, and when  $\varphi \rightarrow \infty$  the variance reduces to the variance of the binomial model, showing that the beta-binomial model is a generalization of the binomial model.

Equation 2 shows how to model the C-to-T damage at a specific base position in the read. We model the position-dependent damage frequency,  $f(x) = k(x)/N(x)$ , see Figure 1, as a function of the distance from the end of the read,  $x$ , with an exponential decay:

$$f(x; A, q, c) = A(1-q)^{x-1} + c.\tag{3}$$

Here  $A$  is the scale factor, or amplitude,  $q$  is the decay rate, and  $c$  is a constant offset, the baseline damage. Since  $x$  is discrete, this is similar to a (modified) geometric sequence starting from  $x = 1$ . The combination of Equation 2 and Equation 3 is illustrated in Figure 2, which shows the position-dependent, decreasing damage

frequency. The figure also shows the increase in uncertainty in the beta-binomial model compared to the binomial-only model.



**Figure 2.** Illustration of the damage model. The figure shows data points as circles and the damage frequency,  $f(x)$ , as a solid line. The amplitude of the damage is  $A$ , the offset is  $c$ , and the relative decrease in damage pr. position is given by  $q$ . The damage uncertainty for a binomial model is shown in dark grey and the uncertainty for a beta-binomial model in light grey.

The damage framework described above is based on the nucleotide misincorporations, i.e. the C-to-T transitions. The background for this data can be from either DNA sequence files mapped to a single genome or from metagenomic data consisting of multiple mapped reads. As such, the damage framework is a general tool for estimating damage based on DNA alignment file(s). In the metagenomic case, metaDMG identifies the lowest common ancestor (LCA) based on the algorithm from the ngsLCA (Wang et al., n.d.).

For each read that maps to multiple reference genomes from separate species, i.e. has multiple alignments, the taxonomic tree is traversed for each alignment until a common ancestor is found. Figure 3 illustrates the LCA for a read that maps to different (sub)species. In this example, the LCA of alignment 1 and 2 is the Subspecies I while the LCA for all four alignments is the Genus X. metaDMG works by default with the NCBI taxonomic database but can also be used with custom databases.

Given the nucleotide misincorporations, either coming from a single-reference alignment file or after LCA in the metagenomic case, eq. (2) and (3) are fitted with a Bayesian model. This is done to ensure the optimal inference of the parameters,  $A$ ,  $q$ , and  $c$ , and to account for the uncertainty in the data. Bayesian inference also allows for the inclusion of domain knowledge in the form of the prior distribution by Bayes theorem. Bayes theorem is based on the law of conditional probability (Barlow, 1993) stating that the probability of two events,  $A$  and  $B$ , both happening,

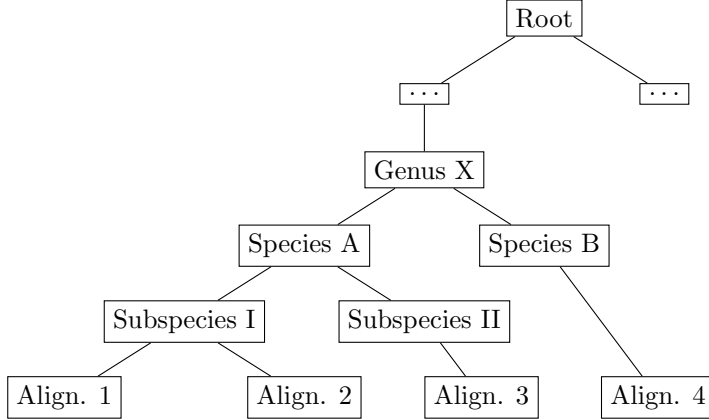
**Figure 3.**

Illustration of the lowest common ancestor (LCA) for taxonomic trees. Here the LCA of alignment 1 and 2 is Subspecies I, while the LCA of all four reads is Genus X. The dots (...) refers to other taxonomic levels, e.g. family and order.

$P(A \cap B)$ , is given by the probability of  $B$ ,  $P(B)$  times the probability of  $A$  given  $B$ ,  $P(A|B)$ :

$$P(A \cap B) = P(B)P(A|B). \quad (4)$$

Similarly,  $P(A \cap B)$  can also be expressed in terms of the probability of  $A$ :

$$P(A \cap B) = P(A)P(B|A). \quad (5)$$

Combining Equation 4 and Equation 5 and rearranging terms gives the Bayes theorem:

$$P(\theta|x) = \frac{P(\theta)P(x|\theta)}{P(x)}, \quad (6)$$

<sup>4</sup> In the case of metaDMG,  $x$  would be the observed deamination frequencies.

with a change of variables where  $x$  now refers to the observed data and  $\theta$  the parameter(s) of the model<sup>4</sup>. The first term in the numerator,  $P(\theta)$ , is the prior distribution and describes the probability distribution assigned to  $\theta$  before observing any data. The second term is the likelihood function,  $P(x|\theta)$ , which is the probability of observing the data,  $x$ , given the parameter(s),  $\theta$ . Together these two terms combine to a compromise between data and prior information.

The numerator,  $P(x)$ , also known as the evidence, can be treated as a data-related normalization factor. In the case of continuous  $\theta$ , this can calculated as the marginalization of the likelihood function over  $\theta$ :

$$P(x) = \int_{\theta} P(x|\theta)P(\theta) d\theta. \quad (7)$$

This equation, however, is often intractable to compute in the higher-dimensional case. Luckily, it can be shown that Markov Chain Monte Carlo (MCMC) sampling can approximate the posterior distribution,  $P(\theta|x)$ , and asymptotically converge to the correct distribution (Gelman, Carlin, et al., 2015).

Traditionally MCMC methods such as Metropolis Hastings (MH) or Gibbs sampling have been used for Bayesian inference, however, these methods are often slow and require a lot of tuning. In the last decades, a new class of MCMC methods have been developed, namely Hamiltonian Monte Carlo (HMC) methods. While traditional MH uses a Gaussian random walk, HMC is a gradient-based MCMC method that uses Hamiltonian dynamics to guide the sampling. This makes HMC more efficient than traditional MCMC methods and allows for sampling from high-dimensional distributions (Betancourt, 2018; Neal, 2011). A particularly efficient variant of HMC is the No-U-Turn Sampler (NUTS). NUTS is a variant of HMC that automatically tunes the step size and number of steps to take in the Hamiltonian dynamics (Homan and Gelman, 2014).

Most statistical domain-specific languages (DSL) such as Stan (Carpenter et al., 2017), Pyro (Bingham et al., 2019), NumPyro (Phan, Pradhan, and Jankowiak, 2019) or Turing.jl (Ge, Xu, and Ghahramani, 2018), implement HMC and in particular the NUTS algorithm. Since `metaDMG` is implemented in Python, NumPyro is used for the Bayesian inference of the damage model, as it is easy to implement and computationally efficient since it which uses JAX (Bradbury et al., 2018) under the hood for automatic differentiation and just-in-time (JIT) compilation.

Even though NumPyro is fast and `metaDMG` is efficiently implemented, the Bayesian inference of the damage model is still computationally expensive. Thus, it was decided to also include a faster, approximate method of Bayesian inference: the maximum a posteriori (MAP) estimate. The MAP estimate is the point estimate of the posterior distribution that maximizes the posterior probability density function, i.e. the posterior mode:

$$\hat{\theta}_{\text{MAP}} = \arg \max_{\theta} P(\theta|x) = \arg \max_{\theta} P(\theta)P(x|\theta), \quad (8)$$

where the second equality is due to the evidence being independent of  $\theta$ . Since this is a point estimate,  $\hat{\theta}_{\text{MAP}}$  does not fully explain the full posterior, however, it is often a good approximation\*. Comparing  $\hat{\theta}_{\text{MAP}}$  to the maximum likelihood estimate (MLE):

$$\hat{\theta}_{\text{MLE}} = \arg \max_{\theta} P(x|\theta), \quad (9)$$

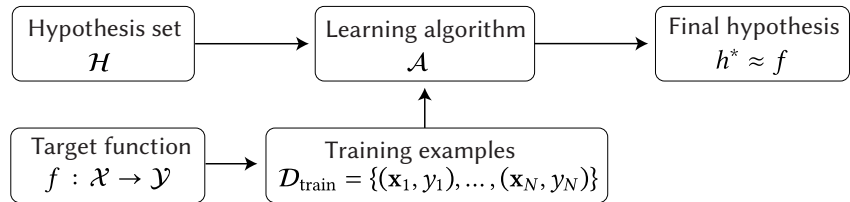
\*Especially when the posterior is unimodal, which it generally is in the case of `metaDMG`.

the MAP estimate can be seen as a regularized version of the MLE estimate (Murphy, 2012). To further optimize the computational efficacy of the MAP estimation in `metaDMG`, the MAP estimation function is JIT compiled using Numba (Lam, Pitrou, and Seibert, 2015) and mathematically optimized with iMinuit (Dembinski et al., 2021).

## 1.2 Anesthesiology – a Machine Learning Approach

This section explains the technical background behind Paper II, see Chapter 3. This study investigates the potential advantages of using a modern machine-learning model compared to classical logistic regression to predict the risk of patients being re-hospitalized after fast-track hip and knee replacements. In particular, the patients were grouped into two groups, where the “risk-patients” all stayed in the hospital for more than four days after the operation or were readmissioned to the hospital within 90 days after the operation. As such, this is a binary classification problem where the patient’s risk-score is predicted based on historical data.

Most classification and regression problems fall under the same machine learning (ML) branch called supervised learning. In supervised learning, the goal is to find the hypothesis  $h^*$  in the hypothesis set  $\mathcal{H}$  that matches the unknown, “true” data-generating function  $f : \mathcal{X} \rightarrow \mathcal{Y}$  optimally, where  $\mathcal{X}$  is the input space and  $\mathcal{Y}$  is the output space. Assuming that we have access to realizations of  $f$ , the so-called training data  $\mathcal{D}_{\text{train}} = \{(\mathbf{x}_i, y_i)\}_{i=1}^N$ , we can use a learning algorithm  $\mathcal{A}$  combined with the training data to estimate  $h^*$  (Abu-Mostafa, Magdon-Ismail, and Lin, 2012). Here  $N$  refers to the number of training samples and  $\mathbf{x}_i$  is the  $i$ th observation with the true label  $y_i$ . This process is illustrated in Figure 4.



**Figure 4.**

Illustration of how to learn from data in a supervised learning setting. Adapted from (Abu-Mostafa, Magdon-Ismail, and Lin, 2012).

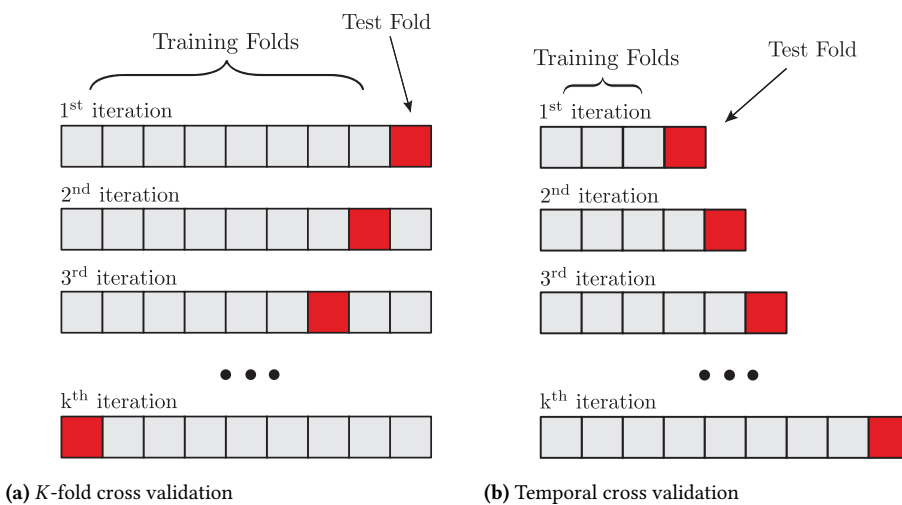
<sup>5</sup> And the hypothesis space thus is significantly larger.

<sup>6</sup> Especially for high cardinality hypothesis sets.

Both the logistic regression (LR) and ML model can be viewed through the lens of Figure 4, just with  $|\mathcal{H}_{\text{LR}}| \ll |\mathcal{H}_{\text{ML}}|$ , i.e. the machine learning model is a lot more complex than the logistic regression model<sup>5</sup>. To predict the performance of  $h^*$  on new, unseen data, the naive method would be to train on all of the data and evaluate on the same, however, this would have a high risk of overfitting the data and thus biasing the predicted performance<sup>6</sup> (Abu-Mostafa, Magdon-Ismail, and Lin, 2012).

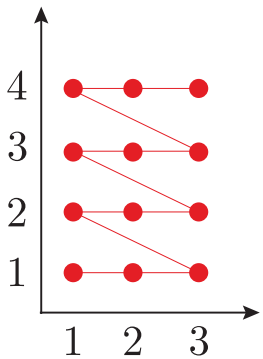
To avoid this and get more accurate estimates of the performance of  $h^*$ , we use a technique called cross-validation (CV). In the simplest way, this can be done by splitting the data into two sets, one called the training and one called the validation set, and then only train on the training set. Afterwards the trained model can be evaluated on the validation set without biasing the performance estimate. This process can further be refined by splitting the data into  $K$  folds and then repeating the process  $K$  times, where each fold is used as the validation set once. This is called  $K$ -fold cross-validation and is illustrated in Figure 5a (Murphy, 2012; Hastie, Tibshirani, and Friedman, 2016).  $K$ -fold cross-validation works well in many cases, yet in the case of temporal data, it also risks introducing bias in the performance estimates, since, in the different folds, it, effectively, is allowed to “look into the future”. The most extreme case of this is shown in the bottom of Figure 5a where the model trains on all future and present data and is then evaluated only on past data. In many time-dependent datasets, this is undesirable. Instead, we use a technique called temporal cross-validation (Tashman, 2000), see Figure 5b, which circumvents this problem by only allowing the model to train on past data and evaluate on future data. As the patient data is time-dependent<sup>7</sup>, this is the technique we use in Paper II.

<sup>7</sup> The fraction of rehospitalizations decreased over time due to surgical improvements.



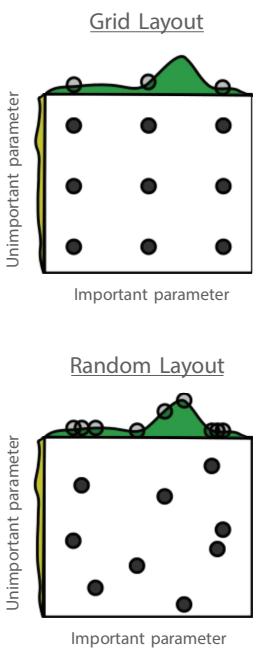
**Figure 5.**  
Two types of cross-validation:  $K$ -fold cross-validation, and temporal cross-validation. Both figures from Michelsen, 2020.

The training of the learning model  $\mathcal{A}$  itself is model-dependent and will not be covered in this thesis, see (Michelsen, 2020) for a more detailed description of the training process. This is not the only way to optimize the performance of  $\mathcal{A}$ , albeit it is the primary one. In addition to the internal parameters of the model, some parameters are external to the model in the sense that they are not optimized by the model itself, but rather by the user. These are called hyperparameters and are often optimized using a technique called hyperparameter optimization (HPO).



**Figure 6.**  
Illustration of grid search.  
Figure from Michelsen,  
2020.

<sup>8</sup> As such, grid search suffers from the curse of dimensionality.



**Figure 7.**  
Illustration comparing grid search to random search.  
The height of green curve is the score-function which has to be optimized. Figure adapted from Bergstra and Bengio, 2012.

In the case of logistic regression, the number of variables to include would be an example of a hyperparameter; in the case of a decision tree model, the depth of the tree. Hyperparameter optimization can be performed in many ways, where the classical one is through grid search, see Figure 6.

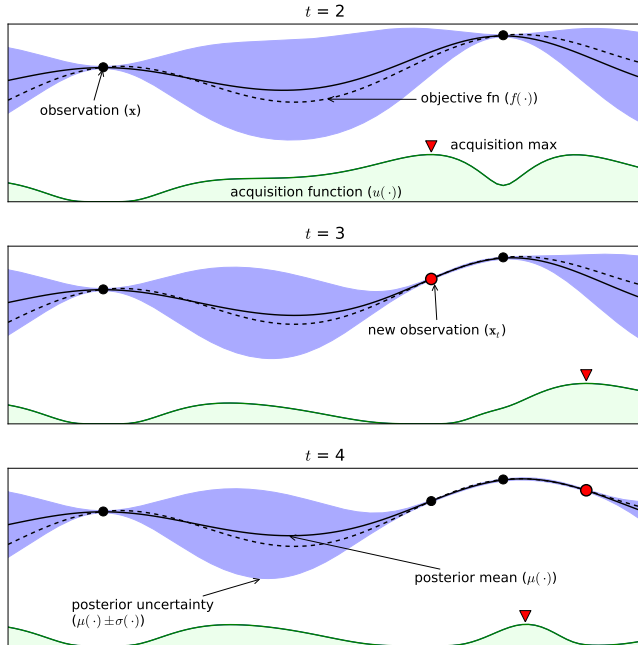
In grid search, all combinations of the hyperparameters (the cartesian product) are tried and the best combination is chosen. This is a simple and intuitive approach, however, it scales exponentially, i.e. very poorly, with the number of hyperparameters<sup>8</sup>. In addition to this, it depends on the user-defined grid, which might not be optimal. To circumvent this, a technique called random search (RS) was developed (Bergstra and Bengio, 2012). Random search is a randomized version of grid search, where the hyperparameters are sampled randomly from a distribution. This allows for a more efficient sampling of the hyperparameter space, see Figure 7, and further lets the user decide on the number of iterations beforehand.

The disadvantage of random search is that all draws are fully independent. While this allows for easy parallelisation of the algorithm, this also means that each new sample might be infinitesimal close in the hyperparameter space to a previous sample with bad performance, which with high probability will thus also have a high loss. An approach that does take the history of the previous samples' performance into consideration is Bayesian optimization (Brochu, Cora, and de Freitas, 2010). In Bayesian optimization each successive hyperparameter is chosen based on an acquisition function, which optimizes the expected improvement in the performance of the model. This is illustrated in Figure 8. This leaves the user with the task of choosing between “exploitation” and “exploration” of the hyperparameter space in the definition of the acquisition function, yet most implementations of bayesian optimization have decent default settings.

We use the Python package Optuna (Akiba et al., 2019) for HPO in Paper IV due to its ease of use and its support for Bayesian optimization. In particular, we use the Tree-structured Parzen Estimator algorithm for the Bayesian optimization and a median stopping rule to minimize optimization time (Bergstra, Bardenet, et al., 2011). This allowed for a good comoprise between optimization time and performance.

While model performance is often paramount, in some fields – such as medicine – being able to explain the model’s predictions is almost as important. This is especially true in the case of medical decision support systems, where the model is used to make decisions about the patient’s treatment. Model explainability helps to build trust in the model, for both the patient and the medical staff alike.

In Paper II, we employ the SHapley Additive exPlanations (SHAP) values which provide estimates on which variables contribute most to the risk score predictions



**Figure 8.** Illustration of the learning process of Bayesian optimization. The previous observations are shown as black dots and the true objective function is shown as a dashed black line. This line is fitted with Gaussian processes which is shown as the solid line with its uncertainty in purple. The acquisition function is shown in green and its maximum decides what the next iteration of the hyper-parameter value(s) should be (Michelsen, 2020).

(Scott M Lundberg and Lee, 2017; Scott M. Lundberg, Erion, et al., 2020). SHAP values allow for not only a global explanation of the model, i.e. which features are most important generally, but also a local explanation, i.e. why a single patient was predicted to be at risk of being re-hospitalized. It has previously been shown that the interaction between SHAP values and medical doctors can improve the performance of anaesthesiologists (Scott M. Lundberg, Nair, et al., 2018).

While the aim of Paper II is to show how modern machine learning techniques can be used to improve the risk prediction process, the usefulness of the SHAP values in a medical context is demonstrated in the paper in Appendix B. The paper uses the SHAP values to compare the preoperative haemoglobin level in the patient with the risk-score, stratified by sex and operation type (knee vs. hip replacement). Currently, the WHO guidelines for the haemoglobin levels are gender specific (Anaemias and Organization, 1968), however, this study finds no significant gender difference and a haemoglobin threshold close to the WHO suggestions for men.

### 1.3 COVID-19 and Agent Based Models

In early 2020, a contagious disease called COVID-19 started to spread in Europe, including Denmark. With new infections showing up faster and faster, governments started to implement different measures to limit the spread of the deadly disease, including lockdowns, travel restrictions, and social distancing, measures

not previously seen in peacetimes since the Spanish flu in 1918. This was the background for the work that we did in 2020 which became the basis for Paper III, see Chapter 4. This paper deals with the development of a new agent based model for COVID-19 in Denmark in collaboration with Statens Serum Institut (SSI), the Danish CDC.

Historically, most mathematical models of infectious diseases were variations of the SIR model, which describe the evolution of a pandemic by approximating all individuals as one population (Kermack, McKendrick, and Walker, 1927). As one of the simplest compartmental models, the susceptible-infectious-recovered (SIR) model is based on a system of three non-linear differential equations that describe the transition between each state, or compartment, of the model (Kröger and Schlickeiser, 2020). Initially the entire population is susceptible until time  $t = 0$  at which some individuals become not only infected, but also infectious, allowing the disease to spread. After having been infectious, the individuals recovers and becomes immune to the disease and are stops being infectious. Several variations of the SIR model exist, including the SIS model, where the recovered individuals become susceptible again (Hethcote, 1989). Another variation is the SEIR model, which includes an exposed state, where individuals are infected but not yet infectious, which is the basis for the model used in Paper III.

SIR-like models suffer from several shortcomings, including the assumptions that the population is homogeneous, and that the disease is transmitted at a constant rate. In reality, neither the population nor the transmission rates are homogenous. These are some of the reasons why we chose to use an agent based model (ABM). Agent based models simulate individual agents in a population that can have complex interactions patterns, e.g. based on their geography (Wilensky and Rand, 2015).

In particular, we implemented a continuous-time, stochastic, spatial ABM using the Gillespie algorithm, a stochastic simulation algorithm (Gillespie, 1977). The model is JIT compiled with Numba (Lam, Pitrou, and Seibert, 2015) to speed up the simulation, allowing simulating the Danish population of 5.8 million people in a couple of hours instead of days. The model allows for the individual tuning of the three main effects; A) heterogeneities in the infection strength<sup>9</sup>, B) number of connections<sup>10</sup>, C) and the spatial clustering of the agents. In the absence of any of these effects, we find that the ABM's predictions matches the SIR model's predictions within  $\pm 5\%$ . Once we allowed for spatial clustering, we found that the epidemic developed faster and with a higher infection peak compared to the SIR model, but that the total number of infected in the end of the epidemic was lower.

In real-life scenarios, one does not have the opportunity to let the epidemic run loose and afterwards evaluate the strength of the epidemic; the goal is to predict the

<sup>9</sup> allowing *super-shedders*

<sup>10</sup> allowing *super-connectors*

intensity in the very beginning of the epidemic and implement lockdown-related measured based on this estimate. In the second part of Paper III, we show that once spatial clustering is introduced, fitting standard SEIR-models to infection numbers from the first few days of the epidemic, predictions are overestimated by a factor of two. The results are a significant over-estimation of the impact of the epidemic. Since the population is highly susceptible in the beginning of an epidemic, this also highlights the benefits of early lockdowns to reduce the effect of the super connectors.

The developed ABM was further used by SSI to estimate the effect of contact tracing related to COVID-19 in Denmark, see Appendix C. It was further used to estimate spread of the “alpha” variant of COVID-19 (B.1.1.7) in Denmark, see Appendix D. Based on data available January 2nd 2021, the model predicted that the “alpha” variant would be the dominant variant in Denmark February 10–20, 2021. It became the dominant variant in Week 7: February 15–21, 2021 (Bager et al., 2021).

#### 1.4 *Diffusion Models and Bayesian Model Comparison*

The similarity between family members and the degree to which siblings resemble one another has long been a mystery in human history. People have always thought about the balance between nature and nurture, as in the famous fairy tale “The Ugly Duckling” by Hans Christian Andersen from 1843. These questions were addressed two decades later, when Gregor Mendel founded genetics as a modern, scientific discipline with his studies on trait inheritance in pea plants (Mendel, Gregor, 1866).

A century later a major breakthrough was when Watson and Crick discovered the double helix structure of DNA (Watson and Crick, 1953). This lead to other important discoveries within genetics, such as the development of DNA sequencing allowing the scientist to identify the genetic makeup for a specific cell. In 2008, the first human genome was sequenced and since then multiple Next-Generation Sequencing methods (NGS) has allowed for cheap, high-quality, in-depth sequencing of genetical samples (Genomics and Mobley, 2021). Since then, the field of genetics has grown exponentially and has become a central part of modern biology today.

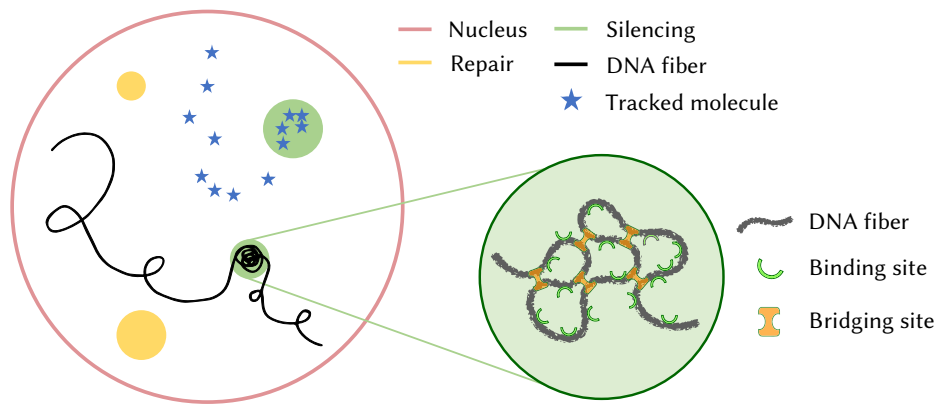
While Section 1.1 discusses the behaviour of ancient DNA, Paper IV focusses on how living cells work and, in particular, how they regulate the transcription of DNA in the cell nucleus. Despite the fact that all cells share DNA, the regulation and expression of the genes stored within can vary. The mechanism of the cell-

specific expression and silencing of specific genomic regions are one of the most fundamental biological challenges.

Currently, different biological models try to explain the physical principles creating the heterogeneous environment in the cell nucleus of eukaryotic cells. One of these is the polymer-bridging model (PBM) that models the micro compartments called the foci. The cell nucleus contains two different types of loci; the repair foci and the silencing foci. Paper IV studies the physical mechanism of the formation of the silencing foci.

Figure 9 illustrates the parts of the cell nucleus relevant to the polymer-bridging model. Inside the nucleus, DNA fibers are curled up and some parts of the DNA locate inside the silencing foci. Inside the silencing foci, the PBM predicts binding and bridging sites that interact with the DNA fiber through the SIR proteins, which is up-regulated inside the the region of the foci (Heltberg et al., 2021). The silent Information Regulator (SIR) proteins repress the underlying genes, and, due to the increased concentration inside the focus, the foci are termed silencing foci.

**Figure 9.**  
Illustration of the cell nucleus. The nucleus membrane is shown in red and the repair foci in yellow. The black line represents the DNA fiber which is curled up in the silencing foci in green. The right side of the figure shows a zoomed in view of the silencing foci according to the polymer-bridging model with the binding and bridging sites that interact with the SIR proteins. The tracking of the SIR proteins is shown as blue stars. Partly adapted from (Heltberg et al., 2021).



With the use of single particle tracking and photoactivated localization microscopy, it is possible to track the individual SIR protein at high temporal and spatial resolution (Oswald et al., 2014; Manley et al., 2008). As the SIR proteins are assumed to follow a diffusion process, the tracking allows for the determination of the diffusion coefficients of cell nucleus, which help quantify the heterogeneous structure in the nucleus.

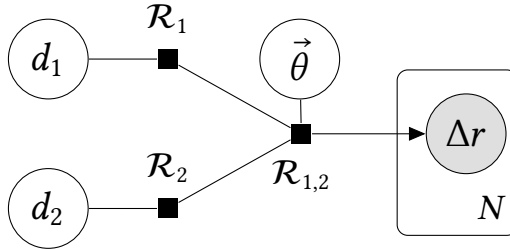
Assuming classical Brownian motion in 2D, the displacement lengths,  $\Delta r_i$ , defined as the distances between subsequent observations  $\vec{x}$ :

$$\Delta r_i = \|\vec{x}_{i+1} - \vec{x}_i\|, \quad (10)$$

follows a Rayleigh distribution:

$$\text{Rayleigh}(r; \sigma) = \frac{r}{\sigma^2} e^{-r^2/(2\sigma^2)} \quad r > 0, \quad (11)$$

with scale parameter  $\sigma = \sqrt{2d\tau}$ , where  $d$  is the diffusion coefficient and  $\tau$  is the time between observations (Anderson et al., 1992). Using Bayesian mixture models, the switch diffusion process is a simple model describing the system, (Baker, 2021). With  $K = 2$  diffusion states, Figure 10 illustrates the model in directed factor graph notation (Dietz, 2022). It shows how the two diffusion coefficients,  $d_1$  and  $d_2$ , each define their own Rayleigh distribution,  $\mathcal{R}_k$ , which are then combined to a mixture distribution,  $\mathcal{R}_{1,2}$ , with mixing probabilities  $\vec{\theta}$ . The measured data,  $\Delta r$ , are  $N$  realisations from this mixture distribution.



The diffusion model illustrated in Figure 10 with  $K = 2$  diffusion states can be extended to  $K$  states, where data shows that both a simpler  $K = 1$  model, the  $K = 2$  model, and a more advanced model with  $K = 3$  diffusion states, all yields appropriate results. Remembering that the formation of the foci depends on the physical properties of the cell nucleus, it is important to be able to evaluate the different models since they provide different diffusion estimates.

The models are compared using the Widely Applicable Information Criterion (WAIC) (Watanabe, 2010) which is a generalized version of the Akaike information criterion (AIC), useful for Bayesian model comparison (Gelman, Hwang, and Vehtari, 2014). The WAIC is an approximation of the out-of-sample loss of the model and is defined as:

$$\text{WAIC} = -2 \left( \underbrace{\text{lppd}}_{\text{accuracy}} - \underbrace{p_{\text{WAIC}}}_{\text{penalty}} \right), \quad (12)$$

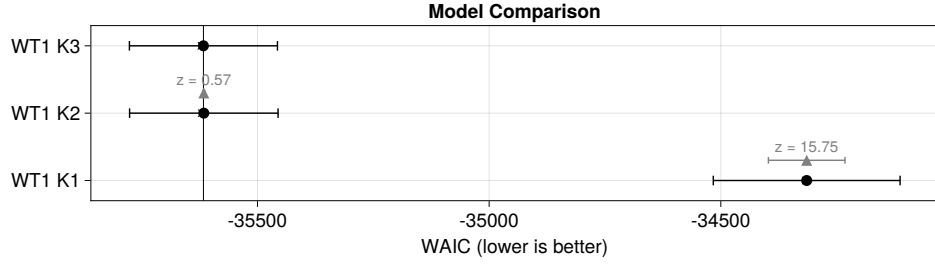
where the log-pointwise-predictive-density (lppd) is a Bayesian version of the accuracy of the model and  $p_{\text{WAIC}}$  is a penalty term that penalizes the model for the effective number of parameters (McElreath, 2020). To compare two models, the model with the lowest WAIC is preferred, however, the difference between the WAICs should also be considered. The results for the WT1 dataset from Paper

**Figure 10.**  
A graphical representation of the Bayesian model case of two diffusion components using the directed factor graph notation (Dietz, 2022). Here  $d_1$  is the diffusion coefficient,  $\mathcal{R}_1$  is the  $d$ -parameterized Rayleigh distribution and  $\mathcal{R}_{1,2}$  is the mixture model of the Rayleigh distributions with a  $\vec{\theta}$  prior.

IV is shown in Figure 11. This figure shows the WAIC in black for the  $K_1$ ,  $K_2$  and  $K_3$  models along with their uncertainties and it is easily seen that the model with only a single diffusion component does not perform well. The difference between the WAIC of the model and the best performing model ( $K_3$ ) is shown in grey,  $\Delta_{A,B}$ , where the  $z$ -value above the error bars are the number of sigmas the difference is from zero:

$$z = \frac{\Delta_{A,B}}{\sigma_{\Delta_{A,B}}}. \quad (13)$$

Following Occam's razor, the  $K_2$  model is chosen as the optimal model, since the difference between the  $K_2$  model and the  $K_3$  model, the best performing one, is statistically non-significant ( $z < 2$ ).



**Figure 11.**

Comparison between diffusion models with  $K = 1$ ,  $K = 2$ , or  $K = 3$  diffusion coefficients for the Wild Type 1 data (WT1). The x-axis shows the WAIC score, where lower values indicate higher-performing models. The WAIC-score for each model is shown in black along with its uncertainty. The difference in WAIC-scores between the model and the best performing model (WT1 K3) is shown in grey with  $z$  being the number of standard deviations between them.

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## **2** *Paper I*

The following pages contain the paper:

**Christian Michelsen**, Mikkel W. Pedersen, Antonio Fernandez-Guerra, Lei Zhao, Troels C. Petersen, Thorfinn S. Korneliussen (2022). “metaDMG: An Ancient DNA Damage Toolkit”.

## metaDMG:

# <sup>2</sup> An Ancient DNA Damage Toolkit

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## Abstract

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**Data availability:** Data is available on [Zenodo](#) or the [Github](#) repository.

**Funding:** This work was supported by Carlsberg Foundation Young Researcher Fellowship awarded by the Carlsberg Foundation [CF19-0712], and the Lundbeck Foundation Centre for Disease Evolution: [R302-2018-2155 to L.Z]. The funders had no role in the decision to publish.

**Competing interests:** The author declare no competing interests.

10. 1. **Motivation** Under favourable conditions DNA molecules can persist for more than two million years (Kjaer et al in press). Such genetic remains make up invaluable resources to study past assemblages, populations and even the evolution of species. However, DNA is subjected to enzymatic, chemical and mechanical degradation, and hence over time decrease to ultra low concentrations which makes it highly prone to contamination by modern sources that are rich in DNA. Strict precautions and criteria(Llamas et al., 2017; Gilbert et al., 2005; Champlot et al., 2010) are therefore necessary to ensure that DNA from modern sources does not appear in the final data and to authenticate that the DNA is ancient. The most generally accepted and widely applied authenticity for ancient DNA studies is to test for elevated deaminated cytosine residues towards the termini of the molecules - DNA damage (Briggs et al., 2007; Dabney, Meyer, and Pääbo, 2013). To date this has primarily been used for single organisms (Jónsson et al., 2013)) and recently for read assemblies (Borry et al., 2021), however these methods have not been designed, nor are they computationally up-scalable for calculating DNA damage for ancient metagenome with tens-hundreds of thousands of species.
24. 2. **Methods** Here we present metaDMG, a novel framework that takes advantage of the information already contained within standard alignment files to compute and statically

evaluate misincorporations due to DNA damage. It thus bypasses any need for initial  
 28 classification, splitting reads by individual organisms, realigning these to the reference genome and lastly parse alignments to mapDamage2.0 (Jónsson et al., 2013). We  
 30 furthermore, implemented a Bayesian approach that combines a geometric damage profile with a beta-binomial model to fit the entire model to the individual misincorporations at all  
 32 taxonomic levels. metaDMG were hereafter benchmarked using simulated data of single genomes, metagenomes and tested on published datasets, before comparing its  
 34 performance with pydamage.

3. **Results** we find metaDMG to be a factor 10 faster than previous methods and more accurate  
 36 even for complex metagenomes with tens of thousands of species. Our simulation show  
 38 that metaDMG can estimate DNA damage at taxonomic levels with less than 100 reads and that uncertainties decreases with increased number of reads, but also that our estimate is more significant with increased C-T misincorporations.  
 40 we also show that sequencing errors does not influence our ability to estimate DNA damage.  
 42 improve the damage estimates in precision and time compared to previous methods.

4. **Conclusion** metaDMG is a state-of-the-art programme suite for computing DNA damage estimation, nucleotide misincorporation, gc-content, DNA fragmentation of simple and complex ancient genomic datasets. Additionally it includes PMDtool statistics (Skoglund et al., 2014) that allow for extraction of reads with damage, making it a complete tools package for ancient DNA damage authentication.

48 **keywords:** ancient DNA, DNA damage estimation, DNA damage, metaDMG, metagenomics, .

## 50 1 | INTRODUCTION

Throughout an organism life it contaminates its environment with DNA, cells or tissue and hereby  
 52 leave genetic traces behind. As the cell leaves its host, DNA repair mechanisms stops and the DNA are subjected to intra and extra-cellular enzymatic, chemical and mechanical degradation, resulting  
 54 in fragmentation and molecule alterations that over time lead to the characteristics of ancient DNA (Briggs et al., 2007; Dabney, Meyer, and Pääbo, 2013). Ancient DNA (aDNA) has been shown to

56 persist in a diverse variety of environmental contexts, e.g. within fossils such as bones, soft-tissue  
and hair, as well as in geological sediments, archaeological layers, ice-cores, permafrost soil for  
58 thousands and even millions of years (Kjaer et al in review), (Cappellini et al., 2018). Common for  
all is that they have an accumulated amount of deaminated cytosines towards the termini of the  
60 DNA strand, which when amplified for sequencing results in mis-incorporations of thymines on the  
cytoses (Dabney, Meyer, and Pääbo, 2013; Ginolhac et al., 2011). This postmortem damage with  
62 regards to DNA is characterized by the four Briggs parameters (Briggs et al., 2007). First, a ancient  
DNA molecule tend to be short, and is likely to have a single stranded overhang due to hydrolytic  
64 cleavage of the  $\beta$ -N-glycosidic bond which results in a nucleotide loss and eventual breakage of the  
DNA. On this overhang the cytosine is especially exposed to deamination which results in the ob-  
66 served high proportion of  $C \rightarrow T$  substitutions at the single stranded overhang  $\epsilon_{ss}$ , and a somewhat  
higher  $C \rightarrow T$  at the double stranded part  $\epsilon_{ds}$ . The length of the single stranded part (*overhang*)  
68 follows a geometric distribution  $\lambda$ , and finally there might be breaks at the backbone in the double  
stranded part  $v$ . It is possible to estimate these four Briggs parameters Jónsson et al., 2013 but  
70 these four parameters are rarely used directly for asserting "ancientness", and researchers work-  
ing with ancient DNA tend to simply use the empirical  $C \rightarrow T$  on the first position of the fragment  
72 together with other supporting summary statistic of the experiment. Estimating mis-incorporation  
due to DNA damage, molecule fragmentation, and nick frequencies have become standard for sin-  
74 gle individual sources like hair, bones, teeth and also applied on small subsets of species in ancient  
environmental metagenomes (Mikkel W. Pedersen et al., 2016; Murchie et al., 2021; Zavala et al.,  
76 2021; Wang, Mikkel Winther Pedersen, et al., 2021). While this is a relatively fast process for single  
individuals it becomes increasingly demanding, iterative and time consuming as the samples and  
78 the diversity within increases, as in the case for metagenomes from ancient soil, sediments, dental  
calculus, coprolites and other ancient environmental sources. It has therefore been practice to es-  
80 timate damage for only the key taxa of interest in a metagenome, as a metagenomic sample easily  
includes tens of thousands of different taxonomic entities, that would make a complete estimate  
82 an impossible task.

To overcome this limitation, we designed a program suite `metaDMG` with novel test statistics that  
84 takes into account all relevant information provided alignments to both single or multiple refer-  
ence genomes. We find `metaDMG`, to estimate DNA damage both faster, more accurate and able to  
86 process complex metagenomes within hours. `metaDMG` is designed for and up-scales equally with

the increasingly large datasets that are currently generated in the field of ancient environmental DNA. However, it also outperforms standard tools that estimate DNA damage in single genomes and samples with low complexity. Furthermore, it can even compute an global damage estimate for a metagenome as a whole. Importantly, metaDMG is compatible with the NCBI taxonomy and use ngsLCA (add Wang et al. 2021 ngsLCA paper) to perform a naïve last common ancestor classification of the aligned reads to get precise damage estimates at all taxonomic nodes. Lastly, it is also designed to accommodate custom taxonomies and hence metagenomic assembled genomes (MAGs) as references.

To test metaDMG specificity, sensitivity and performance we use multiple sets of simulations and test parameters which all show that metaDMG not only outperforms existing methods in the case of single-genome damage estimation but also enables fast and accurate estimation in ancient metagenomes. Lastly, we apply our metaDMG on a representative mix of nine different metagenomes that span the variety of environments hitherto published, including lakes sediments, cave sediments and ancient chewed birch tar.

## 2 | METHODS & MATERIALS

Perhaps the most basic bioinformatic analyses is the difference between two nucleotide sequences. This assumes that we have a haploid representation of our target organisms and larger differences can be interpreted as larger genetic differences. Obtaining a haploid representation is none trivial, firstly our target organism might not be haploid and we need to construct a consensus genome, secondly data from modern day sequencers are essentially a sampling with replacement process and we need to infer the relative location of each of the possible millions or even billions of short DNA fragments, this is the process which is called mapping or alignment. Thirdly, and the focus for this manuscript, is the quantification of the presence of postmortem damage (PMD) in DNA. PMD mainly manifests as an excess of cytosine to thymine substitutions at the termini of fragments that has been prepared for sequencing. A priori we can not directly observe these actual biochemical changes but we can align each fragment and consider the difference between reference and read as possible PMD, and it is even possible to use the excess of C to T at the single fragment level to separate modern from ancient (data with PMD) (Skoglund et al., 2014). Expanding from the single read all reads for a sequencing experiment and genome to tabulate the overall substitution or

116 mismatch rates to obtain a statistic of the damage (Borry et al., 2021) or even estimate the four  
Briggs parameters that is traditionally used to characterize the damage signal (Jónsson et al., 2013).

118 We have devised a general ancient DNA damage toolkit with a special emphasis in a metagenomic setting which implements and expands existing relevant methods but also expands with  
120 several state of the art novel methodologies. At the most basic level we have reimplemented the approach given in (Skoglund et al., 2014) which allows for the extracting and separation of highly  
122 damaged DNA reads. Secondly under the assumption of vast amounts of data we have defined a full multinomial regression model building on the method in (Cabanski et al., 2012), we show that  
124 this will give superior and stable results if it is possible to obtain high depth and coverage data. However, in standard ancient DNA context it is generally not possible to obtain vast amounts of  
126 data and we propose two novel tests statistics that is especially suited for this scenario. To our knowledge there are no currently available methods that is geared towards damage analysis in a  
128 metagenomic setting and existing approaches are essentially based on remapping against the single target organism and does not take into account any possible issues with regards to reads being  
130 well assigned or specified. Our solution called metaDMG (pronounced metadamage), estimates the damage patterns in metagenomic samples in a three step approach. First, the lowest common  
132 ancestor (LCA) for each read (mapped to a multi-species reference database) is computed and the  
the mismatch matrix for each leaf node (e.g. taxonomic ID or contig, depending on the database  
134 used) is computed based on the mapped reads. Second, metaDMG fits a damage model to each leaf node to compute the ancient damage estimates. Finally, the results are visualized in the metaDMG  
136 dashboard, which is a state of the art graphical user interface that allows for fast and user-friendly interaction with the results for further downstream analysis and visualization.

### 138 2.1 | Lowest Common Ancestor and Mismatch matrices

For environmental DNA (eDNA) studies we routinely apply a competitive alignment approach where  
140 we consider all possible alignments for a given read. Each read is mapped against a multi species  
reference databases (e.g. nucleotide or RefSeq from NCBI or custom downloaded). A single read  
142 might map to a highly conserved gene that is shared across higher taxonomic ranks such as class  
or even domains. This read will not provide relevant information due to the generality, whereas a  
144 read that maps solely to a single species or species from a genus would be indicative of the read  
being well classified. We seek to obtain the pattern or signal of damage which is done by the tab-

<sup>146</sup> ulation of the cycle specific mismatch rates between our reference and observed sequence for all  
well classified reads.

<sup>148</sup> In details we compute the lowest common ancestor (lca) for all alignments for each read,  
this is done using (Wang, T. S. Korneliussen, et al., 2022) and if a read is well classified or properly  
<sup>150</sup> assigned based on a user defined threshold (species, genus or family) we tabulate the mismatches  
for each cycle, if a read is not well assigned it is discarded. Pending on the run mode we allow  
<sup>152</sup> for the construction of these mismatch tables on three different levels. Either we obtain a basic  
single global mismatch matrix, which could be relevant in a standard single genome aDNA study  
<sup>154</sup> and similar to the tabulation used in (Jónsson et al., 2013). Secondly we can obtain per reference  
counts or if a taxonomy database has been supplied we allow for the aggregation from leaf nodes  
<sup>156</sup> to the internal taxonomic ranks towards the root.

To suit as many users as possible, metaDMG takes as input an alignment file (.bam, .sam, or  
<sup>158</sup> .sam.gz), where each read is hereafter allowed an equal chance to map against the multiple references. One read can therefore attract multiple alignments, and we thus first seek to find the lowest  
<sup>160</sup> common ancestor (LCA) among the alignments based on the tree structure from the databases and  
a user defined read-reference similarity interval (Wang, T. S. Korneliussen, et al., 2022). Note that  
<sup>162</sup> metaDMG is not limited to the NCBI database and allow for custom databases as well.

Regardless of runmode or weighing scheme used in the possible aggregation we obtain the  
<sup>164</sup> nucleotide substitution frequencies across reads which provides us with the position dependent  
mismatch matrices,  $\underline{\underline{M}}(x)$ , with  $x$  denoting the position in the read, starting from 1. At a specific  
<sup>166</sup> position,  $M_{ref \rightarrow obs}(x)$  describes the number of nucleotides that was mapped to a reference base  $B_{ref}$   
but observed to be  $B_{obs}$ , where  $B \in \{A, C, G, T\}$ . The number of C to T transitions, e.g., is denoted  
<sup>168</sup> as  $M_{C \rightarrow T}(x)$ .

When calculating the mismatch matrix, two different approaches can be taken. Either all align-  
<sup>170</sup> ments of the read will be counted, which we will refer to as weight-type 0, or the counts will be  
normalized by the number of alignments of each read; weight-type 1 (default).

## <sup>172</sup> 2.2 | Damage Estimation

The damage pattern observed in aDNA has several features which are well characterized. By mod-  
<sup>174</sup> elling these, one can construct observables sensitive to aDNA signal. We model the damage pat-  
terns seen in ancient DNA by looking exclusively at the  $C \rightarrow T$  transitions in the forward direction

<sup>176</sup> (5') and the  $G \rightarrow A$  transitions in the reverse direction (3'). For each LCA, we denote the number of transitions  $k(x)$  as:

$$\begin{aligned} \text{178} \quad k(x) = & \begin{cases} M_{C \rightarrow T}(x) & \text{for } x > 0 \quad (\text{forward}) \\ M_{G \rightarrow A}(x) & \text{for } x < 0 \quad (\text{reverse}) \end{cases} \end{aligned} \quad (1)$$

<sup>180</sup> and the number of the reference counts  $N(x)$ :

$$\begin{aligned} \text{182} \quad N(x) = & \begin{cases} \sum_{i \in B} M_{C \rightarrow i}(x) & \text{for } x > 0 \quad (\text{forward}) \\ \sum_{i \in B} M_{G \rightarrow i}(x) & \text{for } x < 0 \quad (\text{reverse}), \end{cases} \end{aligned} \quad (2)$$

<sup>184</sup> where the sum is over all four bases. The damage frequency is thus  $f(x) = k(x)/N(x)$ . A natural choice of likelihood model would be the binomial distribution. However, we found that a binomial likelihood lacks the flexibility needed to deal with the large amount of variance (overdispersion) <sup>186</sup> we found in the data due to bad references and misalignments.

To accommodate overdispersion, we instead apply a beta-binomial distribution,  $\mathcal{P}_{\text{BetaBinomial}}$ , which

<sup>1</sup> Note that we do not parameterize the beta distribution in terms of the common  $(\alpha, \beta)$  parameterization, but instead using the more intuitive  $(\mu, \phi)$  parameterization. One can re-parameterize  $(\alpha, \beta) \rightarrow (\mu, \phi)$  using the following two equations:  $\mu = \frac{\alpha}{\alpha + \beta}$  and  $\phi = \alpha + \beta$  (<sup>188</sup> Cepeda-Cuervo and Cifuentes-Amado, 2017).

<sup>188</sup> treats the probability,  $p$ , as a random variable following a beta distribution<sup>1</sup> with mean  $\mu$  and concentration  $\phi$ :  $p \sim \text{Beta}(\mu, \phi)$ . The beta-binomial distribution has the the following probability density function:

$$\mathcal{P}_{\text{BetaBinomial}}(k | N, \mu, \phi) = \binom{N}{k} \frac{B(k + \mu\phi, N - k + \phi(1 - \mu))}{B(\mu\phi, \phi(1 - \mu))}, \quad (3)$$

where  $B$  is defined as the beta function:

$$B(x, y) = \frac{\Gamma(x)\Gamma(y)}{\Gamma(x+y)}, \quad (4)$$

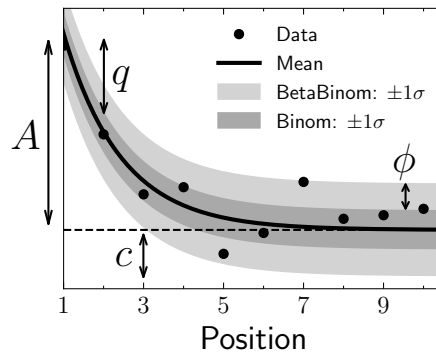
<sup>196</sup> with  $\Gamma(\cdot)$  being the gamma function (Cepeda-Cuervo and Cifuentes-Amado, 2017).

The close resemblance to a binomial model is most easily seen by comparing the mean and <sup>198</sup> variance of a random variable  $k$  following a beta-binomial distribution,  $k \sim \mathcal{P}_{\text{BetaBinomial}}(N, \mu, \phi)$ :

$$\begin{aligned} \mathbb{E}[k] &= N\mu \\ \text{200} \quad \mathbb{V}[k] &= N\mu(1 - \mu) \frac{\phi + N}{\phi + 1}. \end{aligned} \quad (5)$$

<sup>202</sup> The expected value of  $k$  is similar to that of a binomial distribution and the variance of the beta-binomial distribution reduces to a binomial distribution as  $\phi \rightarrow \infty$ . The beta-binomial distribution can thus be seen as a generalization of the binomial distribution.

<sup>204</sup> Note that both equation (3) and (5) relates to damage at a specific base position, i.e. for a single  $k$  and  $N$ . To estimate the overall damage in the entire read using the position dependent counts,



**Figure 1.** Illustration of the damage model. The figure shows data points as circles and the damage,  $f(x)$ , as a solid line. The amplitude of the damage is  $A$ , the offset is  $c$ , and the relative decrease in damage pr. position is given by  $q$ . The damage uncertainty for a binomial model is shown in dark grey and the uncertainty for a beta-binomial model in light grey. The additional uncertainty of the beta-binomial model, compared to the binomial model, is related to  $\phi$ , see equation (5).

$k(x)$  and  $N(x)$ , we model  $\mu$  as position dependent,  $\mu(x)$ , and assume a position-independent concentration,  $\phi$ . We model the damage frequency with a modified geometric sequence, i.e. exponential decreasing for discrete values of  $x$ :

$$\tilde{f}(x; A, q, c) = A(1 - q)^{|x|-1} + c. \quad (6)$$

Here  $A$  is the amplitude of the damage and  $q$  is the relative decrease of damage pr. position. A background,  $c$ , was added to reflect the fact that the mismatch between the read and reference might be due to other factors than just ancient damage. As such, we allow for a non-zero amount of damage, even as  $x \rightarrow \infty$ . This is visualized in Fig. 1 along with a comparison between the classical binomial model and the beta-binomial model.

To estimate the fit parameters,  $A$ ,  $q$ ,  $c$ , and  $\phi$ , we apply Bayesian inference to utilize domain specific knowledge in the form of priors. We assume weakly informative beta-priors<sup>2</sup> for both  $A$ ,  $q$ , and  $c$ . In addition to this, we assume an exponential prior on  $\phi$  with the requirement of  $\phi > 2$  to

<sup>2</sup> Parameterized as  $(\mu, \phi)$

<sup>218</sup> avoid too much focus on 0-or-1 probabilities (McElreath, 2020). The final model is thus:

$$\begin{aligned}
 \text{[}A\text{ prior]} & A \sim \text{Beta}(0.1, 10) \\
 \text{[}q\text{ prior]} & q \sim \text{Beta}(0.2, 5) \\
 \text{[}c\text{ prior]} & c \sim \text{Beta}(0.1, 10) \\
 \text{[}\phi\text{ prior]} & \phi \sim 2 + \text{Exponential}(1000) \\
 \text{[likelihood]} & k_i \sim \mathcal{P}_{\text{BetaBinomial}}(N_i, \tilde{f}(x_i; A, q, c), \phi),
 \end{aligned} \tag{7}$$

where  $i$  is an index running over all positions.

<sup>226</sup> We define the damage due to deamination,  $D$ , as the background-subtracted damage frequency at the first position:  $D \equiv \tilde{f}(|x| = 1) - c$ . As such,  $D$  is the damage related to ancientness. Using the <sup>228</sup> properties of the beta-binomial distribution, eq. (5), we find the mean and variance of the damage:

$$\begin{aligned}
 \mathbb{E}[D] & \equiv \bar{D} = A \\
 \mathbb{V}[D] & \equiv \sigma_D^2 = \frac{A(1-A)}{N} \frac{\phi+N}{\phi+1}.
 \end{aligned} \tag{8}$$

<sup>230</sup> Since  $D$  estimates the overexpression of damage due to ancientness, not only the mean is relevant but also the certainty of  $D > 0$ . We quantify this through the significance  $Z = \bar{D}/\sigma_D$  <sup>232</sup> which is thus the number of standard deviations ("sigmas") away from zero. Assuming a Gaussian distribution of  $D$ ,  $Z > 2$  would indicate a probability of  $D$  being larger than zero, i.e. containing <sup>234</sup> ancient damage, with more than 97.7% probability. These two values allows us to not only quantify the amount of ancient damage (ie.  $\bar{D}$ ) but also the certainty of this damage ( $Z$ ) without even having <sup>236</sup> to run multiple models and comparing these.

We perform the Bayesian inference of the parameters models using Hamiltonian Monte Carlo <sup>238</sup> (HMC) sampling which is a particular of Monte Carlo Markov Chain (MCMC) algorithm (Betancourt, <sup>240</sup> 2018). Specifically, we use the NUTS implementation in NumPyro (Phan, Pradhan, and Jankowiak, <sup>242</sup> 2019), a Python package which uses JAX (Bradbury et al., 2018) under the hood for automatic differentiation and JIT compilation. We treat each leaf node of the LCA as being independent and generate 1000 MCMC samples after an initial 500 samples as warm up.

Since running the full Bayesian model is computationally expensive, we also allow for a faster, <sup>244</sup> approximate method by just fitting the maximum a posteriori probability (MAP) estimate. We use iMinuit (Dembinski et al., 2021) for the MAP optimization with Numba acceleration (Lam, Pitrou, <sup>246</sup> and Seibert, 2015) for even faster run times. On a Macbook M1 Pro model from 2021, the timings

for running the full Bayesian model is  $1.41 \pm 0.04$  s/fit and for the MAP it is  $4.34 \pm 0.07$  ms/fit, showing  
 248 more than a 2 order increase in performance (around 300x) for the approximate model. Both  
 models allow for easy parallelisation to decrease the computation time.

### 250 2.3 | Visualisation

We provide an interactive dashboard to properly visualise the results from the modelling phase,  
 252 see <https://metadmg.onrender.com/> for an example. The dashboard allows for filtering, styling and  
 variable selection, visualizing the mismatch matrix related to a specific leaf node, and exporting of  
 254 both fit results and plots. By filtering, we include both filtering by sample, by specific cuts in the fit  
 results (e.g. requiring  $D$  to be above a certain threshold), and even by taxonomic level (e.g. only  
 256 looking tax IDs that are part of the Mammalia class). We greatly believe that a visual overview of  
 the fit results increase understanding of the data at hand. The dashboard is implemented with  
 258 Plotly plots and incorporated into a Dash dashboard (Plotly, 2015).

## 3 | SIMULATION STUDY

260 To ascertain the performance of our test statistic and implementation we performed various rig-  
 orous simulation studies to quantify possible issues with bias and accuracy in a synthetic setting  
 262 that should mimick the various issues and complications that exist with real world data. We con-  
 ducted two sets of simulations, one to gauge the performance of the damage model itself and one  
 264 to determine the performance of the full metaDMG pipeline, i.e. both LCA and damage model.

### 3.1 | Single-genome Simulations

266 The first set of simulations was performed by taking a single, representative genome and adding  
 post mortem damage together with sequencing noise. This was followed by a standard mapping  
 268 step and finally damage estimation using metaDMG. The deamination was applied using NGSNGS  
 (Henriksen, Zhao, and T. Korneliussen, 2022) which is a recent implementation of the original  
 270 Briggs model similar to Gargammel (Neukamm, Peltzer, and Nieselt, 2021). In this step we vary  
 the simulated amount of damage added (in particular the single-stranded DNA deamination,  $\delta_{ss}$ )  
 272 in the original Briggs model (Briggs et al., 2007)), the number of reads, and the fragment length  
 distribution.

274       `./ngsngs [something]`

bowtie2 [something]

276 We chose five different, representative genomes, in each of these varying the three simulation  
 parameters. These genomes were the homo sapiens, the betula, and three microbial organisms  
 278 with respectively low, median, and high amount of GC-content. For each of these simulations, we  
 performed 100 independent replicates to measure the variability of the parameter estimation and  
 280 quantify the robustness of the estimates. We simulated eight different sets of damage (approx-  
 imately 0%, 1%, 2%, 5%, 10%, 15%, 20%, 30%), 13 sets of different number of reads (10, 25, 50, 100, 250,  
 282 500, 1.000, 2.500, 5.000, 10.000, 25.000, 50.000, 100.000), three sets of different fragment length distri-  
 butions (samples from a *log-normal* distribution with mean 35, 60, and 90, each with a standard  
 284 deviation of 10), and five different genomes, each simulation set repeated 100 times.

In addition to this, we also create 1000 repetitions of the non-damaged simulations for Homo  
 286 Sapiens to be able to gauge the risk of finding false positives. Finally, to show that the damage esti-  
 mates that metaDMG provides are independent of the contig size, we artificially create three different  
 288 genomes by sampling 1.000, 10.000 or 100.000 different basepairs from a uniform categorical dis-  
 tribution of {A, C, G, T}.

290 To be able to compare our estimates to a known value, we generate 1.000.000 reads using  
 NGSNGS without any added sequencing noise for each of other sets of simulation parameters.  
 292 The difference in damage frequency at position 1 and 15 is then the value to compare to:

$$D_{\text{known}} = \frac{f(x=1) - f(x=15)}{2} + \frac{f(x=-1) - f(x=-15)}{2}, \quad (9)$$

294 where we take the average of the C to T damage frequency difference and the G to A damage  
 frequency difference.

296 The fastq files were simulated with NGSNGS using the above mentioned simulation parameters,  
 all with the same quality scores profiles as used in ART (Huang et al., 2012), based on the Illumina  
 298 HiSeq 2500 (150 bp). The mapping was performed using Bowtie-2 with the –no-unal flag (Langmead  
 and Salzberg, 2012).

### 300 3.2 | Metagenomic Simulations

While the previously mentioned simulation study is perfectly aimed at quantifying the performance  
 302 of the damage model in the case of single-reference genomics it does lack the complexity related  
 to metagenomic samples. Therefore, we also conduct a more advanced simulation study to deter-

304 mine the accuracy of the full metaDMG pipeline.

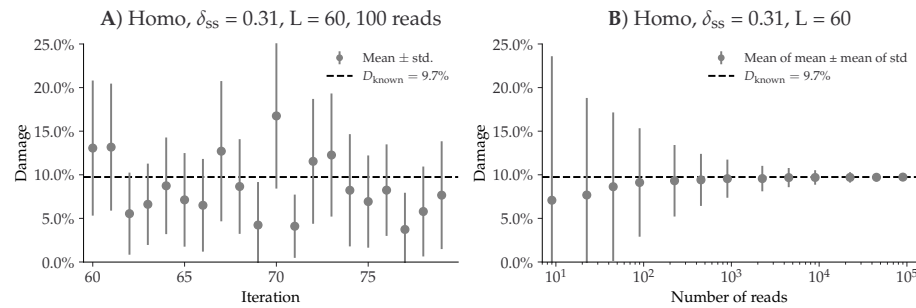
305 The previously mentioned simulation study quantifies the damage model's performance for  
 306 single-reference genomics, but it does not address the complexity of metagenomic studies. There-  
 307 fore, we also conducted a more advanced simulation study to determine the performance of the  
 308 metaDMG pipeline in a standard eDNA setting. Based on an ancient metagenome scenario, we cre-  
 309 ated a synthetic dataset that mimics the composition, fragment length distribution, and damage  
 310 patterns for each genome. We selected X metagenomes (Supp table XXX) covering several environ-  
 311 mental conditions and ages. First, we mapped the reads of each metagenome with bowtie2 against  
 312 a database that contained the GTDB r202 (Parks et al., 2018) species cluster reference sequences,  
 313 all organelles from NCBI RefSeq (NCBI Resource Coordinators, 2018), and the reference sequences  
 314 from CheckV (Nayfach et al., 2021). We used bam-filter 1.0.11 with the flag --read-length-freqs to  
 315 get the mapped read length distribution for each genome and their abundance. The genomes with  
 316 an observed-to-expected coverage ratio greater than 0.75 were kept. The filtered BAM files were  
 317 processed by metaDMG to obtain the misincorporation matrices. The abundance tables, fragment  
 318 length distribution, and misincorporation matrices were used in aMGSIM-smk v0.0.1 (Fernandez-  
 319 Guerra, 2022), a Snakemake workflow (Mölder et al., 2021) that facilitates the generation of many  
 320 synthetic ancient metagenomes. The data used and generated by the workflow can be obtained  
 321 from Figshare link (XXX). We then performed taxonomic profiling using the same parameters used  
 322 for the synthetic reads generated by aMGSIM-smk.

## 4 | RESULTS

323 The accuracy of all methods in metaDMG was tested in various simulation scenarios. In general we  
 324 find that metaDMG yields accurate, precise damage estimates even in extreme low-coverage data.

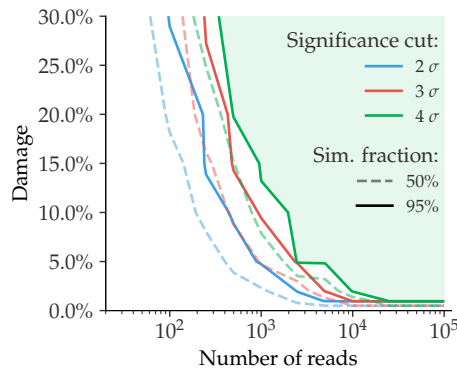
### 326 4.1 | Single-genome Simulations

327 The results of the single-genome simulations can be seen in Figure 2. The left part of the figure  
 328 shows metaDMG damage estimates based on the homo sapiens genome with the Briggs parameter  
 $\delta_{SS} = 0.31$  and a fragment length distribution with mean 60, each of the simulations generated with  
 329 100 simulated reads for 10 representative simulations. When the damage estimates are low, the  
 330 distribution of  $D$  is highly skewed (restricted to positive values) leading to errorbars sometimes  
 331 going into negative damage, which of course represents un-physical values. The right hand side of

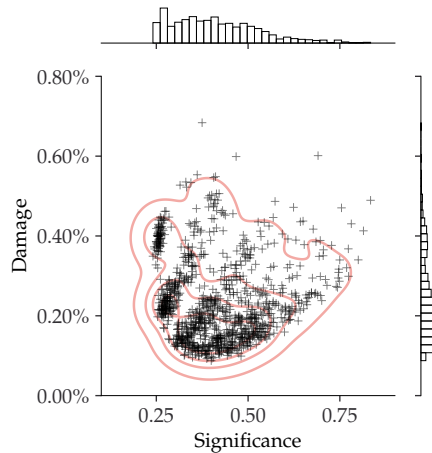


**Figure 2.** Overview of the single-genome simulations based on the homo sapiens genome with the Briggs parameter  $\delta_{SS} = 0.065$  and a fragment length distribution with mean 60. **A)** This plot shows the estimated damage ( $D$ ) of 10 simulations with 100 simulated reads. The grey points show the mean damage (with its standard deviation as errorbars). The known damage ( $D_{known}$ ) is shown as a dashed line, see eq. (9). **B)** This plot shows the average damage as a function of the number of reads. The grey points show the average of the individual means (with the average of the standard deviations as errors).

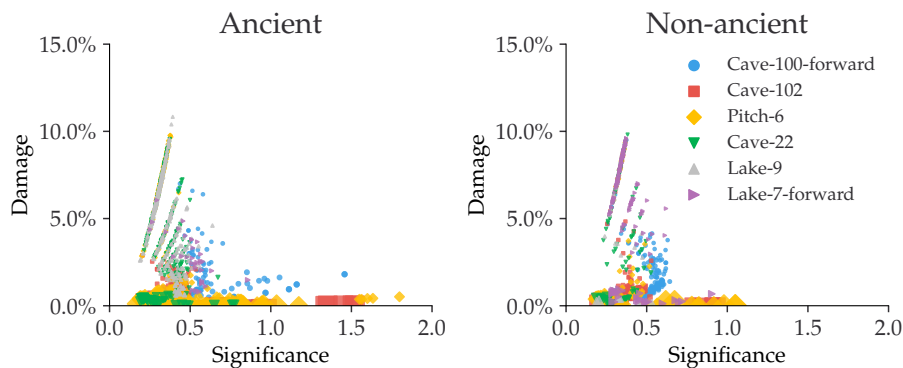
the figure visualizes the average amount of damage across a varying number of reads. This shows  
 344 that the damage estimates converge to the known value with more data, and that one needs more than 100 reads to even get strictly positive damage estimates (when including uncertainties).  
 346 Across more than 5 different species, 3 different fragment length distributions, and 3 different contig length distributions, each with 100 simulations for 104 different sets of simulation parameters, the only difference we note in the damage estimates is between species with low, median, and high GC-levels. In general, species with higher GC-levels exhibit lower variations in their damage  
 348 estimates compared to species with lower GC-levels, leading to high-GC species requiring fewer reads to establish damage estimates.  
 342 Based on the single-genome simulations, we can compute the relationship between the amount of damage in a species and the number of reads required to correctly infer that the given species  
 344 is damaged, see Figure 3. If we want to find damage with a significance of more than 2 (solid blue line) in a sample with around 5% expected damage, it requires about 1000 reads to be 95% certain  
 346 that we will find results this good. Said in other words: given 100 different fits, each with 1000 reads and around 5% damage, one would expect to find damage (with a  $Z > 2$ ) in 95 of the total  
 348 100 samples, on average. If we loose the requirement such that it is okay to only find it in every second fit, it would be enough with only around 250 reads in each fit (dashed blue line).  
 350 Finally, to quantify the risk of incorrectly assigning damage to a non-damaged species, we cre-



**Figure 3.** Given a specific significance cut, the solid contour line shows the relationship between the amount of damage and the number of reads required to be able to correctly infer damage in 95% of the species. The dashed line shows the similar value for a simulation fraction of 50%. The green part of the figure shows the “good” region of number of reads and estimated damage, given than one wants to be more than 95% certain of correctly identifying damage with more than  $4\sigma$  confidence.



**Figure 4.** This figure shows the inferred damage estimates of 1000 independent simulations, each with 1000 reads and no artificial ancient damage applied, with the inferred damage shown on the y-axis and the significance on the x-axis. Each simulation is shown as a single cross and the red lines show the kernel density estimate (KDE) of the damage estimates. The marginal distributions are shown as histograms next to the scatter plot.

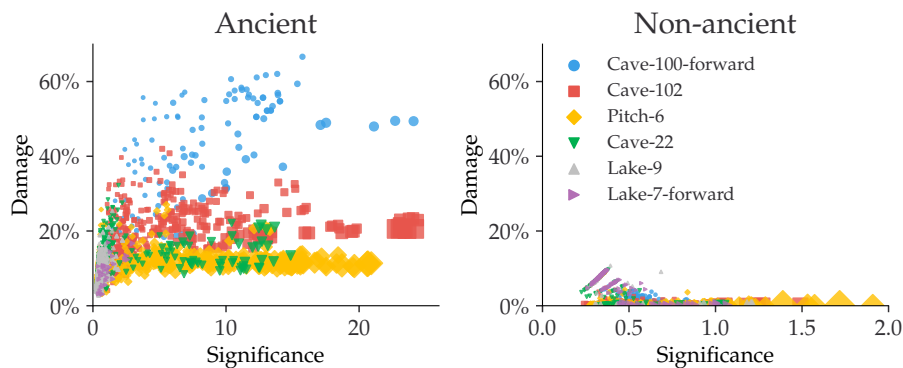


**Figure 5.** Estimated amount of damage as a function of significance using the fragSim data. The left figure shows the damage of the species that we simulated to be ancient (however with no deamination added yet) and the right figure shows the same for the species that are not going to have deamination added.

ated 1000 independent simulations for a varying number of reads, where none of them had any ar-  
 362 tificial ancient damage applied, only sequencing noise. Figure 4 shows the damage ( $D$ ) as a function  
 of the significance ( $Z$ ) for the case of 1000 simulated reads. Even though the estimated damage is  
 364 larger than zero, the damage is non-significant since the significance is less than one. When looking  
 at all the figures across the different number of reads, see Figure bayesian\_zero\_damage\_plots.pdf,  
 366 we note that a loose cut requiring that  $D > 1\%$  and  $Z > 2$  would filter out all of non-damaged  
 points. Overall the conclusion being that our devised test statistic is conservative and has low  
 368 false positive rate.

#### 4.2 | Metagenomic Simulations

360 With the full metagenomic simulation pipeline we can further probe the performance of metaDMG.  
 By looking at the six different metagenomic scenarios at different steps in the pipeline we are able  
 362 to show that metaDMG provides relevant, accurate damage estimates. First of all, we run metaDMG on  
 the six samples after fragmentation with FragSim. Since no deamination has yet been added at  
 364 this step in the pipeline, this is also a test of the risk of getting false positives. The results can be  
 seen in Figure 5 where we see the damage estimates for both the species that we simulate to be  
 366 ancient and the species that we do not add deamination to. We see that the damage estimates are  
 quite similar, as expected, and that our previously established loose cut of  $D > 1\%$  and  $Z > 2$  still  
 368 filters out all of non-damaged points.



**Figure 6.** Estimated amount of damage as a function of significance using the ART data. The left figure shows the damage of the species that we simulated to be ancient and the right figure shows the same for the species that have not had deamination added.

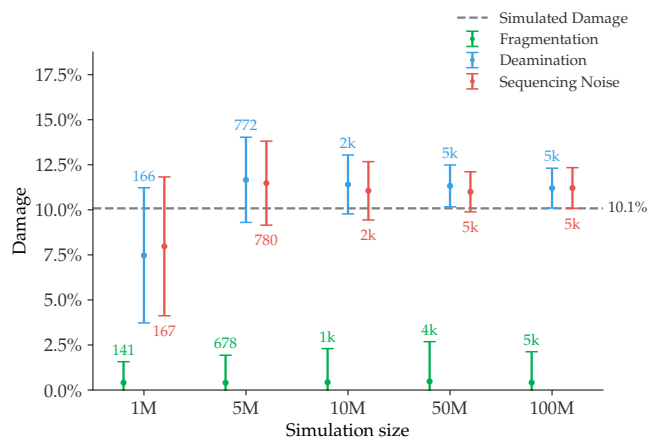
In comparison we can look at Figure 6 which shows the same plot, but after the deamination 370 (deamSim) and sequencing errors (ART) has been added. Here we see a clear difference between the ancient and the non-ancient ones, as expected. The non-ancient species would still not pass 372 the loose cut, however, we note that a large number of the ancient samples would. By looking at Figure 6 we see that not all of the samples show similar amount of damage. These observations 374 are summarised in Table 1 where we see that Cave-100-forward, Cave-102, Pitch-6 all have more than 60% of their ancient species labelled as damaged according to the loose cut, Cave-22 (18%) 376 and Lake-7-forward (12%) a bit lower, while Lake-9 (0.5%) does not show any clear signs of damage. However, once we condition on the requirement of having more than 100 reads, the fraction of 378 ancient species correctly identified as ancient increases to more than 90% for most the samples.

To better understand the damage estimates, we can look at them individually. Figure 7 shows 380 the *Stenotrophomonas maltophilia* species from the Pitch-6 sample. We see that none of the fragmentation-only files were estimated to have damage and that most of the deamination and 382 final files including sequencing errors have damage – at a simulation size of 1 million, the significance of both are  $Z \approx 1.9$ , so this one of the few fits with more than 100 reads that does not 384 pass the loose cut. Furthermore, we notice that the error bars decrease with simulation size, as expected.

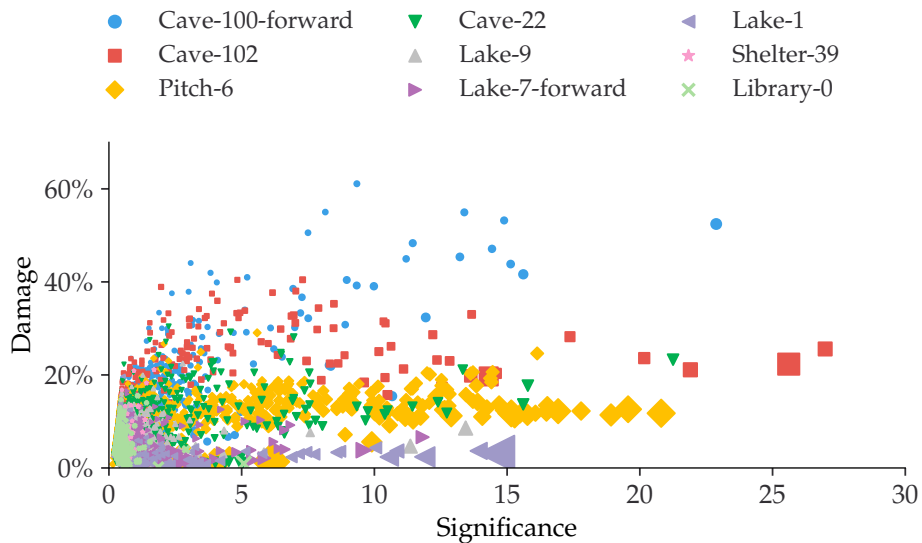
386 The rest of the metagenomic simulation results are shown in Figure XXX.

**Table 1.** Number of ancient species for each of the six simulated samples. The first column is the total number of species, the second column is the total number of species that would pass the loose cut of  $D > 1\%$  and  $Z > 2$ , the third column is the number of species with more than 100 reads, and the final column is the number of species with more than 100 reads that also do pass the cut.

Sample	Total	Pass	+100 Reads	+100 Reads and Pass
Cave-100-forward	135	107	79.3%	88
Cave-102	500	326	65.2%	309
Pitch-6	415	260	62.7%	274
Cave-22	393	71	18.1%	73
Lake-9	410	2	0.5%	8
Lake-7-forward	32	4	12.5%	6



**Figure 7.** Damage estimates of the *Stenotrophomonas maltophilia* species from the Pitch-6 sample. Damage is shown as a function of simulation size, with the fragmentation files in green, the deamination files in blue and the final files including sequencing errors in red. All errors are  $1\sigma$  error bars (standard deviation). The number of reads for each fit is shown as text and since this was a species simulated to have ancient damage, the simulated amount of damage is shown as a dashed grey line.



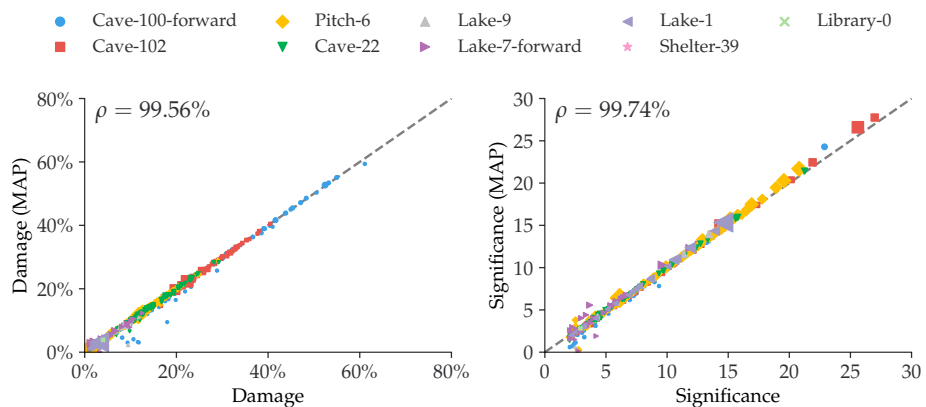
**Figure 8.** Estimated amount of damage as a function of significance using the real data.

#### 4.3 | Real Data

388 The results from running the full metaDMG pipeline on real data can be seen in Figure 8. The figures  
 389 shows Blablabla, real life data here. We find that the loose cut ( $D > 1\%$ ,  $Z > 2$ ) accepts only one of  
 390 the fits from the control test Library-0, which would not have been accepted by more conservative  
 391 cut ( $D > 2\%$ ,  $Z > 3$ , more than 100 reads).

#### 4.4 | Bayesian vs. MAP

392 Due to increased computational burden of running the full Bayesian model compared to faster,  
 393 approximate MAP model, in samples with several thousand species, the MAP model is often the  
 394 most realistic model to use due to time constraints. In this case, it is of course important to know  
 395 that the damage estimates are indeed trustworthy. Figure 9 compares the estimated damage  
 396 between the Bayesian model and the MAP model and the estimated significances for species with  
 397  $D > 1\%$ ,  $Z > 2$  and more than 100 reads. The figure shows that the vast majority of species map  
 398 1:1 between the Bayesian and the MAP model. One should note, though, that the few species with  
 399 the highest mismatch, all are based on forward-only fits, i.e. with no information from the reverse  
 400 strand, which thus leads to less data to base the fits on. For the comparison with no cuts, see  
 401 Figure 1 in appendix.



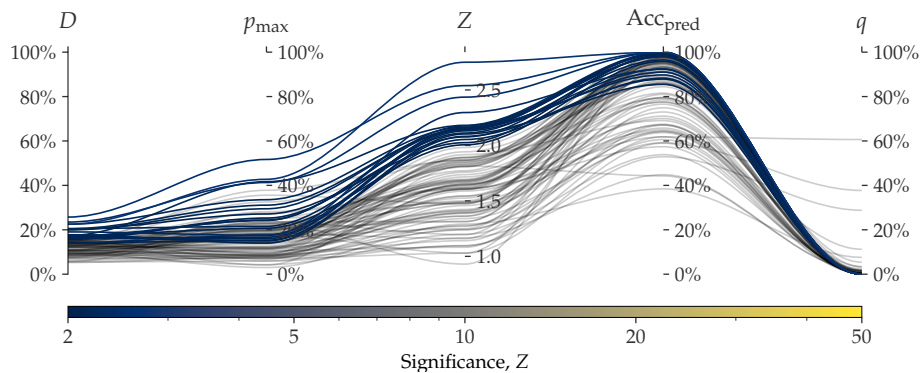
**Figure 9.** Comparison between the full Bayesian model and the fast, approximate, MAP model for the estimated damage and significance. The figure shows data after a loose cut of  $D > 1\%$ ,  $Z > 2$  and more than 100 reads. The dashed, grey line shows the 1:1 ratio and the correlation,  $\rho$ , is shown in the upper right corner.

#### 4.5 | Existing Methods

404 We have also compared `metaDMG` to existing methods such as PyDamage (Borry et al., 2021). Since  
 405 PyDamage does not include the LCA step, this comparison is based on the non-LCA mode (local-  
 406 mode) of `metaDMG`. This mode iterates through the different assigned species for all mapped reads  
 407 and estimates the damage for each. In general, we find that `metaDMG` is more conservative, accurate  
 408 and precise in its damage estimates.

On example of this is can be found in Figure 10, which shows both the `metaDMG` and PyDamage  
 410 results of the 100 *Homo Sapiens* single-genome simulations with 100 reads and 15% added artificial  
 411 damage (and a fragment length distribution with mean 60).

412 To compare the computational performance, we use the Pitch-6 sample which has 11.433  
 413 unique taxa. When using only a single core, PyDamage took 1105 s to compute all fits, while `metaDMG`  
 414 took 88 s, a factor of 12.6x faster. Out of the 88 s, `metaDMG` spent 53 s on the actual fits, the rest was for  
 415 loading and reading the alignment file and computing the mismatch matrices. This makes `metaDMG`  
 416 more than 20x faster than PyDamage for the fit computation. For the rest of the timings, see Table  
 417 2. PyDamage requires the alignment file to be sorted by chromosome position and be supplied  
 418 with an index file, allowing it to iterate fast through the alignment file, at the expense of computa-  
 419 tional load before running the actual damage estimation. `metaDMG` on the other hand requires the  
 420 reads to be sorted by name to minimize the time it takes to run the LCA, which however, is not



**Figure 10.** Parallel Coordinates plot comparing metaDMG and PyDamage for the Homo Sapiens single-genome simulation with 100 reads and 15% added artificial damage. The different axis shows the five different variables: metaDMG-damage ( $D$ , by metaDMG), PyDamage-damage ( $p_{\max}$ , by PyDamage), significance ( $Z$ , by metaDMG), predicted accuracy ( $\text{Acc}_{\text{pred}}$ , by PyDamage), and the p-value ( $q$ , by PyDamage). Each of the 100 simulations are plotted as single lines showing the values of the different dimensions. Simulations with  $D > 1\%$  and  $Z > 2$ , i.e. damaged according to the loose metaDMG cut, are shown in color proportional to their significance. Non-damaged simulations are shown in semi-transparent black lines.

tested in this comparison.

## 422 5 | DISCUSSION

Preliminary work indicates that the computational performance of the models can be even fur-  
424 ther optimized by using Julia (Bezanson et al., 2017), which shows around 7x optimization for the Bayesian model (~ 0.2 s/fit) and 4x for the MAP model (~ 1.1 ms/fit).

- 426     • contig length
- simulation setup
- 428     • more covariates, readlength? other substitutions
- no linkage
- 430     • taxa independence,
- weight
- 432     • improved fishing (pmdtools)
- relevant testcases?
- 434     • basis for environmental deme differences?

**Table 2.** Computational performance of PyDamage and metaDMG. The table contains the times it takes to run either PyDamage or metaDMG on the full Pitch-6 sample containing 11.433 species. The timings are shown for both single-processing case (1 core) and multi-processing (2 and 4 cores). The timings were performed on a Macbook M1 Pro model from 2021. “12.6x” means that metaDMG was 12.6 times faster than PyDamage for that particular test.

Cores	Pitch-6		Pydamage		metaDMG	
	Total	Fits	Total	Fits	Total	Fits
1	1105 s	1102 s	88 s	12.6x	53 s	20.8x
2	592 s	590 s	66 s	9.0x	25 s	23.6x
4	398 s	397 s	54 s	7.4x	14s	28.4x

## 5.1 | Acknowledgment

<sup>436</sup> Acknowledgements here

## 5.2 | Data availability

<sup>438</sup> Source code is hosted at GitHub: <https://github.com/metaDMG-dev>. Sequencing data can be found at: <https://somewhere.com> XXX.

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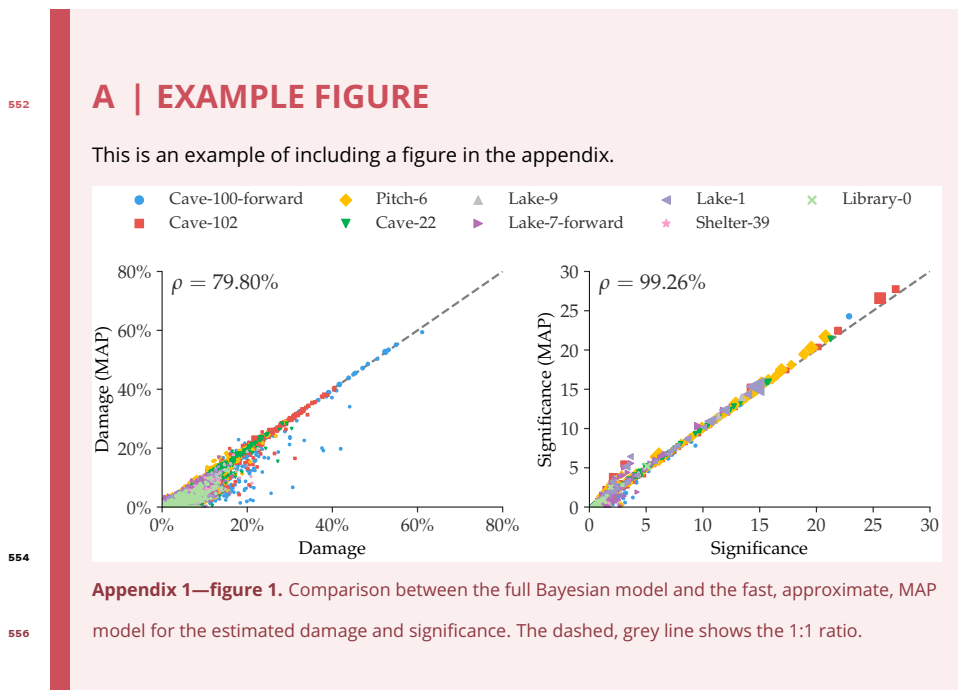
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- 550

## Appendix 1



## 558 Appendix 2

## 560 B | EXAMPLE TABLE

561 This is an example of including a table in the appendix.

562 Appendix 2—table 1. An example table.

Speed (mph)	Driver	Car	Engine	Date
407.447	Craig Breedlove	Spirit of America	GE J47	8/5/63
413.199	Tom Green	Wingfoot Express	WE J46	10/2/64
434.22	Art Arfons	Green Monster	GE J79	10/5/64
468.719	Craig Breedlove	Spirit of America	GE J79	10/13/64
526.277	Craig Breedlove	Spirit of America	GE J79	10/15/65
536.712	Art Arfons	Green Monster	GE J79	10/27/65
555.127	Craig Breedlove	Spirit of America, Sonic 1	GE J79	11/2/65
576.553	Art Arfons	Green Monster	GE J79	11/7/65
600.601	Craig Breedlove	Spirit of America, Sonic 1	GE J79	11/15/65
622.407	Gary Gabelich	Blue Flame	Rocket	10/23/70
633.468	Richard Noble	Thrust 2	RR RG 146	10/4/83
763.035	Andy Green	Thrust SSC	RR Spey	10/15/97

564 Appendix 3

## C | MULTINOMIAL LOGISTIC REGRESSIONS

### 566 Full Multinomial Logistic Regression models

568 Postmortem damages will have impacts on the NGS (next generation sequencing) reads.

570 A common phenomenon is the calling error rates increases from nucleotide C to T due to the cytosine deamination process. Unawareness of this will lead to inaccurate inferences.

572 Evidences show that the magnitude of such changes are related to the positions the site is within a read (the fraction of the ancient DNA). Here we present 3 slightly different ways to unveil the relationship between the calling error rates and the mismatching reference/read pairs as well as the site positions within a read. The methods are based on the multinomial logistic regressions.

### 574 Data Description

576 We perform the regression based on the summary statistic of the mismatch matrix, i.e.,  $\underline{\underline{M}}(x)$ , which is a table which contains the counts of reads of different reference/read categories (in total 16) and positions on the forward/reversed strand (15 positions on each direction).

578 Table 1 and Table 2 give an example of the data format we use for the inference.

Ref.	Read Counts								
	Read	A				C			
		A	C	G	T	A	C	G	T
1	12794053	8325	28769	16073	10404	8045811	8020	2092619	
2	13480290	6812	21107	12102	9151	8260185	6531	1145605	
3	12760253	6131	18859	10327	7772	8385423	5899	914709	
4	12995572	5240	17671	8940	7880	8345892	5252	767237	
5	12930102	4601	17021	8188	8374	8474964	5161	703283	
6	12879355	4684	16435	7536	8726	8571141	4811	643607	
7	12684349	4557	15298	7394	8835	8727254	4762	586674	
8	12585563	4454	15497	7236	8898	8888173	5058	527691	
9	12468622	4309	14704	6942	8948	9076851	4673	481170	
10	12491183	4437	14567	6912	9103	9237982	4702	443329	
11	12430899	4296	14083	6515	9313	9364121	4609	404431	
12	12419506	4226	13985	6503	9342	9357468	4367	371475	
13	12469412	4147	13851	6375	9586	9386737	4588	345390	
14	12549936	4045	13650	6246	9673	9324488	4628	322294	
15	12566555	4174	13499	6213	9735	9305820	4518	301360	
-1	11599167	8800	16164	14851	90888	9613102	10843	19810	
-2	11985637	8769	14044	12040	28799	9561124	7184	18424	
-3	12941743	7805	13861	12001	24988	9400151	6368	15466	
-4	12808985	7141	12885	9889	23067	9509723	5421	14901	
-5	12869585	6954	12100	9428	22349	9464831	5789	13987	
-6	12784911	6440	12080	8735	20556	9566794	6544	14021	
-7	12878349	5946	12311	8225	19480	9566359	6478	16419	
-8	12719722	9521	12156	8131	19226	9725468	6709	23434	
-9	12652860	5634	11940	7671	18035	9762224	6321	31667	
-10	12566817	5448	11850	7178	17353	9701382	6306	37831	
-11	12702498	5309	12092	7568	16121	9526031	6035	43215	
-12	12731940	5207	11933	6856	15637	9533858	5557	47650	
-13	12697647	4989	12199	7153	15072	9508117	5434	51614	
-14	12689924	4944	11891	6816	15050	9525285	5237	bioRxiv preprint doi: https://doi.org/10.1101/555982; this version posted April 29, 2022. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under a CC-BY-NC-ND 4.0 International license.	
	12660634	4746	11753	6732	14815	9561359	5184	59633	

582 **Appendix 3—table 1.** The read counts per position given the reference nucleotides are A or C of an  
ancient human data. The negative position indices are the position on the reversed strand. In the  
manuscript, the elements (the values of a specific nucleotide read counts per position given the  
reference nucleotide is A or C) in this table are denoted as  $M_{A \rightarrow i}(x)$  or  $M_{C \rightarrow i}(x)$ .

584

Read Counts

Ref.	G									T										
	A			C			G			T			A			C			G	
Read	A	C	G	T	A	C	G	T	A	C	G	T	A	C	G	T	A	C	G	T
1	16389	8976	9639767	86584	11733	15878	8351	11718463												
2	17614	6483	9510149	26655	10761	13958	7011	11974947												
3	15164	5949	9488917	23374	9509	13767	6046	12839015												
4	14844	5186	9566468	21960	8170	12509	5585	12721790												
5	14005	5612	9497118	20468	7186	11991	5233	12795244												
6	13671	6195	9622572	19096	6948	11683	4790	12686645												
7	16648	6394	9609855	18594	6203	12122	4780	12794172												
8	23659	6405	9768666	17341	6131	11847	4758	12626614												
9	31680	6139	9785449	17034	5998	12040	4469	12579260												
10	38484	5982	9700857	16235	5487	11546	4175	12513653												
11	44665	5722	9536341	15284	5651	12044	4176	12646627												
12	48949	5371	9547134	14569	5449	11663	4060	12684645												
13	53076	5234	9543953	14090	5262	11785	4046	12631297												
14	57343	5186	9551477	13855	5257	11768	4006	12624840												
15	61236	5137	9583481	13667	5122	11733	3947	12612416												
-1	2078554	7947	8096447	11847	15732	28461	8551	12890628												
-2	1138478	6656	8232666	10760	12299	20759	6999	13446882												
-3	921712	5970	8399013	8643	10514	18226	6564	12718084												
-4	775038	5720	8319235	8416	9415	17800	5388	12977322												
-5	710955	5499	8462058	8926	8526	17088	4911	12886576												
-6	647761	5052	8545455	9193	7640	16351	4879	12852322												
-7	593854	4872	8693834	9318	7600	15523	5048	12664576												
-8	535542	7828	8889921	9399	7163	18704	4718	12510123												
-9	486549	4696	9075263	9522	7109	14547	4611	12409220												
-10	448895	4622	9226758	9432	6816	14567	4668	12438344												
-11	409027	4654	9352528	9544	6575	14019	4611	12388650												
-12	376069	4637	9344701	9419	6511	13874	4486	12390148												
-13	350609	4655	9384853	9885	6197	13877	4327	12432024												
-14	326760	4595	9337266	9889	5986	13928	4403	b124909901												
-15	305014	4541	9310617	10065	5919	13442	4232	12529684												

**Appendix 3—table 2.** The read counts per position given the reference nucleotides are G or T of the same human data as in Table 1. The negative position indices are the position on the reversed strand. In the manuscript, the elements (the values of a specific nucleotide read counts per position given the reference nucleotide is G or T) in this table are denoted as  $M_{G \rightarrow i}(x)$  or  $M_{T \rightarrow i}(x)$ .

The terminology used here might not be standard. The term full regression here is to distinguish itself from the folded regression discussed later, which simply means inferring the coefficients of forward strand and reversed strand separately. Full regression includes both unconditional regression and conditional regression. The unconditional regression's objective is to infer the probability of observing a read of nucleotide  $j$  and its reference is  $i$  at position  $x$ , i.e.,  $P_{i \rightarrow j}(x)$  while the conditional regression's target is to estimate the probability of observing a read of nucleotide  $j$  given its reference is  $i$  at position  $x$ , i.e.,  $P_{j|i}(x)$ . Their relationship is as follows:

$$P_{j|i}(x) = \frac{P_{i \rightarrow j}(x)}{\sum_{j \in B} P_{i \rightarrow j}(x)}.$$

So in fact, unconditional regression can give us more detailed inferred results (extra information the nucleotide composition per position of the reference, which may be related to the prepared libraries).

#### Unconditional Regression Likelihood

The unconditional regression's log-likelihood function is defined as follows,

$$\begin{aligned} l_1 &= \sum_x \sum_{i,j \in B} M_{i \rightarrow j}(x) \log P_{i \rightarrow j}(x) \\ &= \sum_x \left[ M(x) \log P_{T \rightarrow T}(x) + \sum_{(i,j) \neq (T,T)} M_{i \rightarrow j}(x) \log \frac{P_{i \rightarrow j}(x)}{P_{T \rightarrow T}(x)} \right], \end{aligned} \quad (10)$$

where  $M(x) = \sum_{i,j \in B} M_{i \rightarrow j}(x)$ . According to the multinomial logistic regression, we assume,

$$\log \frac{P_{i \rightarrow j}(x)}{P_{T \rightarrow T}(x)} = \sum_{n=0}^{\text{order}} \alpha_{i,j,x,n} x^n \quad (11)$$

Applying Equation 11 to Equation 10, we have

$$l_1 = \sum_x \left\{ -M(x) \log \left[ 1 + \sum_{(i,j) \neq (T,T)} \exp \left( \sum_{n=0}^{\text{order}} \alpha_{i,j,x,n} x^n \right) \right] + \sum_{(i,j) \neq (T,T)} M_{i \rightarrow j}(x) \sum_{n=0}^{\text{order}} \alpha_{i,j,x,n} x^n \right\} \quad (12)$$

620 The number of inferred parameters ( $\alpha_{i,j,x,n}$ ), for the full conditional regression is  $30 \times (\text{order} + 1)$ .

622 And the relevant derivatives of the unconditional regression likelihood are as follows,

$$\frac{\partial l_1}{\partial \alpha_{i,j,x,n}} = -M(x) \frac{x^n \exp\left(\sum_{n=0}^{\text{order}} \alpha_{i,j,x,n} x^n\right)}{1 + \sum_{(i,j) \neq (T,T)} \exp\left(\sum_{n=0}^{\text{order}} \alpha_{i,j,x,n} x^n\right)} + M_{i \rightarrow j}(x)x^n. \quad (13)$$

### 626 Conditional Regression Likelihood

628 Viewed as the sum of log-likelihoods given the reference nucleotide  $i \in B$ , the conditional  
regression's log-likelihood function is,

$$\begin{aligned} l_2 &= \sum_{i \in B} \sum_x \sum_{j \in B} M_{i \rightarrow j}(x) \log P_{j|i}(x) \\ &= \sum_{i \in B} \sum_x \left[ M_i(x) \log P_{T|i}(x) + \sum_{j \neq T} M_{i \rightarrow j}(x) \log \frac{P_{j|i}(x)}{P_{T|i}(x)} \right], \end{aligned} \quad (14)$$

632 where  $M_i(x) = \sum_{j \in B} M_{i \rightarrow j}(x)$ . Furthermore, if we assume,

$$\log \frac{P_{j|i}(x)}{P_{T|i}(x)} = \sum_{n=0}^{\text{order}} \beta_{i,j,x,n} x^n \quad (15)$$

634 By applying Equation 15 to Equation 14, we can obtain,

$$l_2 = \sum_{i \in B} \sum_x \left\{ -M_i(x) \log \left[ 1 + \sum_{j \neq T} \exp \left( \sum_{n=0}^{\text{order}} \beta_{i,j,x,n} x^n \right) \right] + \sum_{j \neq T} M_{i \rightarrow j}(x) \sum_{n=0}^{\text{order}} \beta_{i,j,x,n} x^n \right\} \quad (16)$$

636 The number of inferred parameters ( $\beta_{i,j,x,n}$ ) for the full unconditional regression is  $24 \times$   
638 (order + 1). And the relevant derivatives of the conditional likelihood are as follows,

$$\frac{\partial l_2}{\partial \beta_{i,j,x,n}} = -M_i(x) \frac{x^n \exp\left(\sum_{n=0}^{\text{order}} \beta_{i,j,x,n} x^n\right)}{1 + \sum_{j \neq T} \exp\left(\sum_{n=0}^{\text{order}} \beta_{i,j,x,n} x^n\right)} + M_{i \rightarrow j}(x)x^n. \quad (17)$$

### Folded Regression

The folded regressions use the same log-likelihood functions as the full regression (i.e., Equation 12 and 16) but are conducted based on a presumable symmetric PMD pattern, i.e., the probability of  $C \rightarrow T$  at the position  $x$  of an random chosen ancient DNA strand is assumed to equal to the probability of  $G \rightarrow A$  at the position  $-x$ . Such an theoretical assumption go match the current ancient library preparation process [Meyer's paper and

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Rasmus H's paper].

650

$$\alpha_{i,j,x,n} = \alpha_{c(i),c(j),-x,n}, \quad (18)$$

652

$$\beta_{i,j,x,n} = \beta_{c(i),c(j),-x,n}, \quad (19)$$

654

where  $c(i)$  means the complimentary nucleotide of the nucleotide  $i$ , e.g.,  $c(A) = T$  and  $c(G) = C$ .

656

By doing the folded regression, we halve the number of inferred parameters ( $\alpha_{i,j,x,n}$  or  $\beta_{i,j,x,n}$ ). Hence The number of inferred parameters for the folded unconditional regression is  $15 \times (\text{order} + 1)$ , and that of folded conditional regression is  $12 \times (\text{order} + 1)$ .

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### Results for multinomial logistic regression

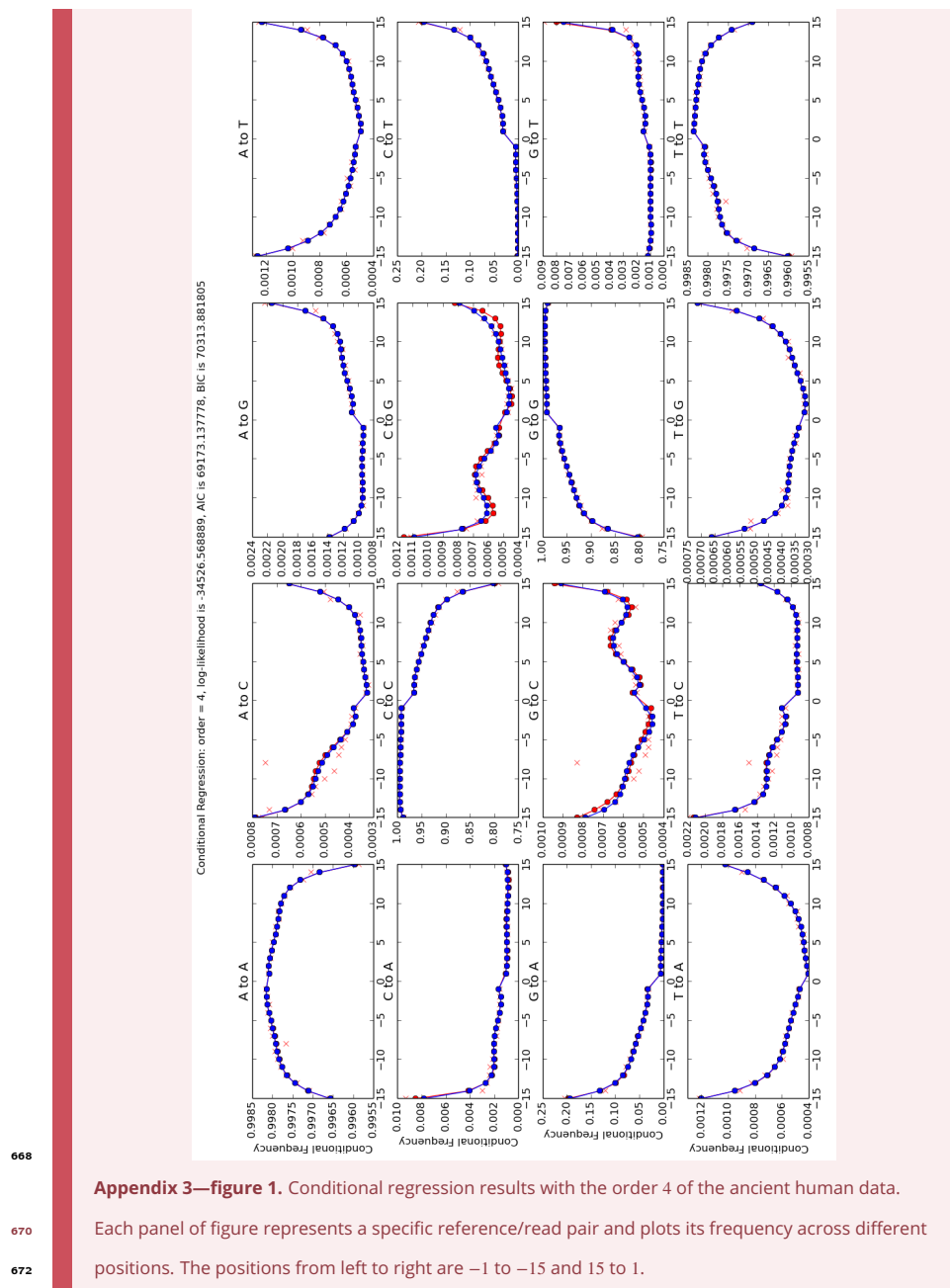
660

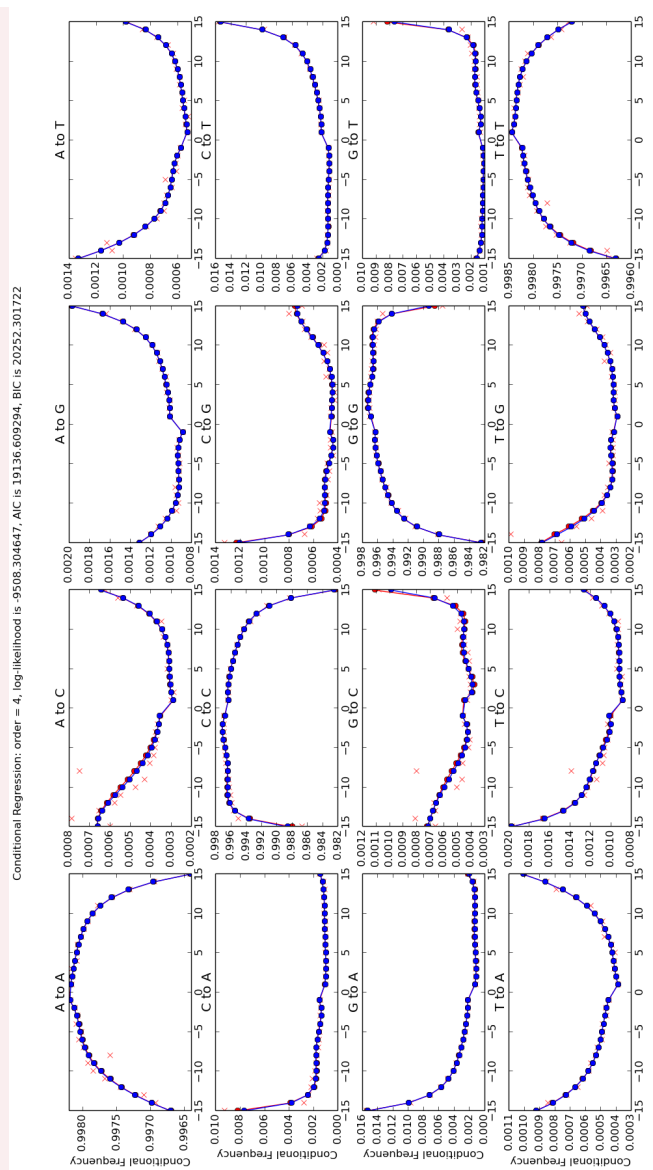
Currently, the optimization of the likelihood functions are based on the C++ library of gsl and use the function `gsl_multimin_fminimizer_nmsimplex2`. with the initial searching point is set to be the results of logistic regression. We here present here 4 figures pertaining to showcase the performance of our model. The regression methods are based on the summary statistic of the counts of mismatches and the optimization is therefore in the scale of miliseconds. Fig. 1 and Fig. 2 are the conditional regression results of the ancient and control human data correspondingly. And Fig. 3 and Fig. 4 are the folded conditional regression results of the same data as above. Our codes can also do the unconditional regression, but I have not generated the results for now.

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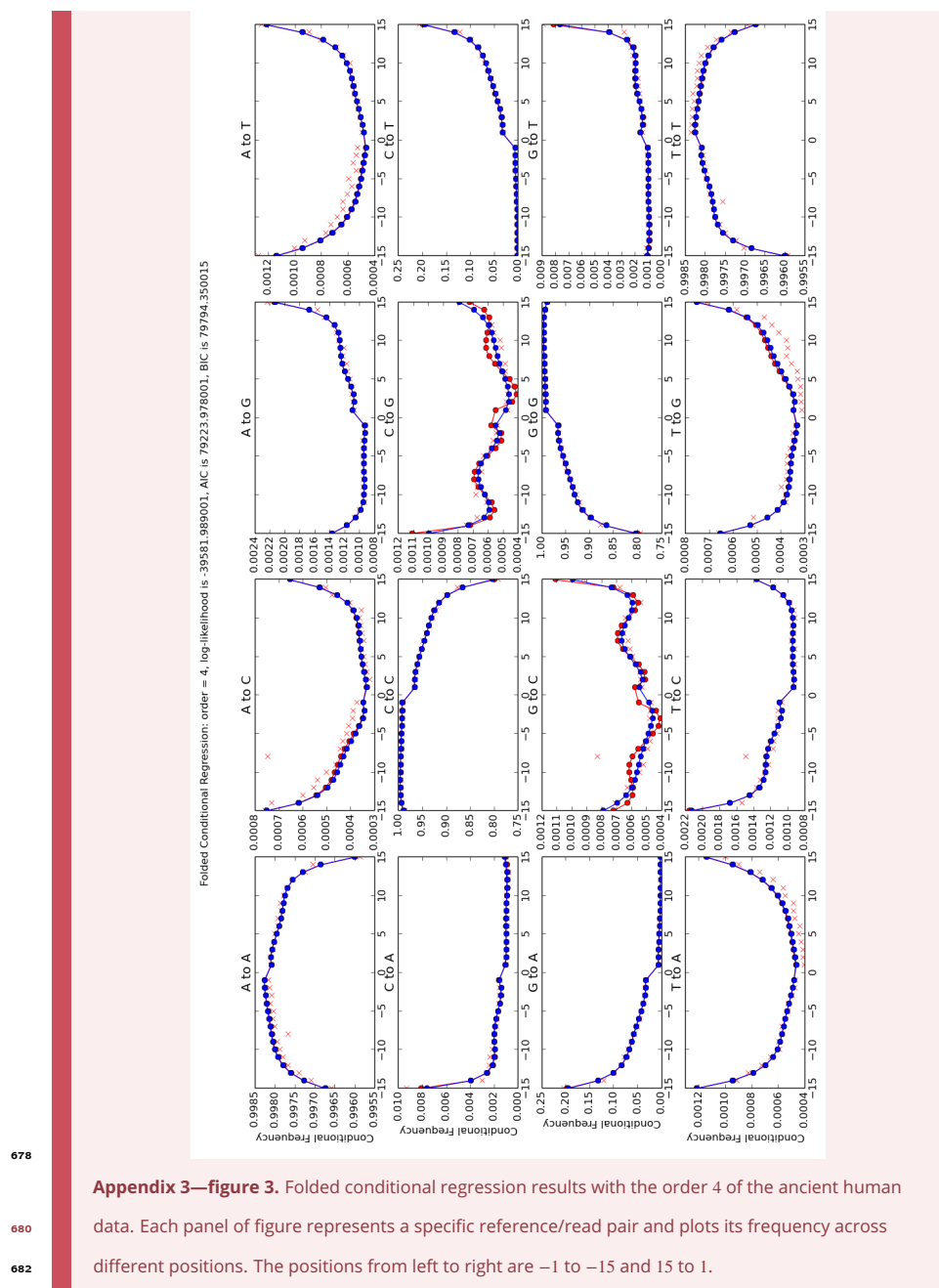


674

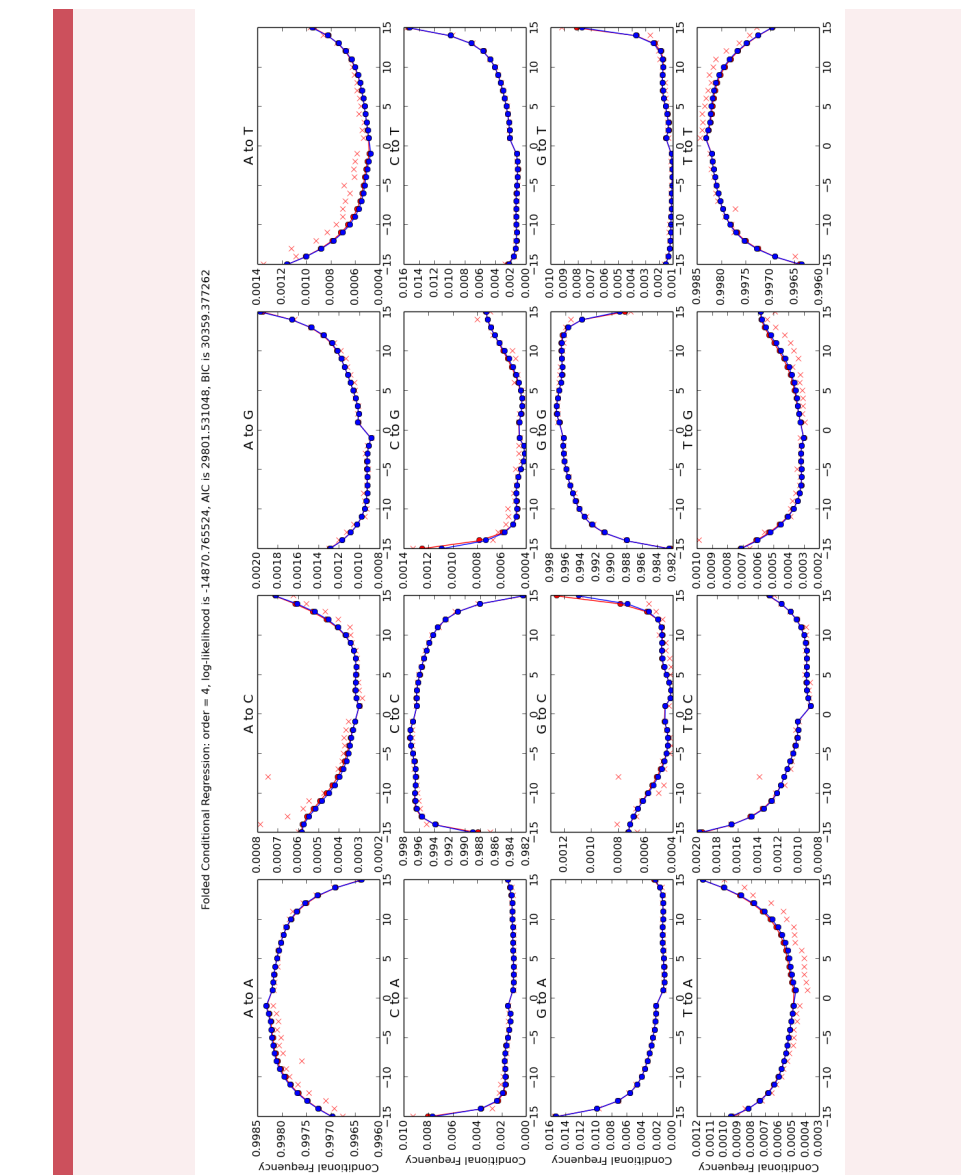
**Appendix 3—figure 2.** Conditional regression results with the order 4 of the control human data.

Each panel of figure represents a specific reference/read pair and plots its frequency across different positions. The positions from left to right are  $-1$  to  $-15$  and  $15$  to  $1$ .

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**Appendix 3—figure 3.** Folded conditional regression results with the order 4 of the ancient human data. Each panel of figure represents a specific reference/read pair and plots its frequency across different positions. The positions from left to right are  $-1$  to  $-15$  and  $15$  to  $1$ .



684  
686  
**Appendix 3—figure 4.** Folded conditional regression results with the order 4 of the control human data. Each panel of figure represents a specific reference/read pair and plots its frequency across different positions. The positions from left to right are  $-1$  to  $-15$  and  $15$  to  $1$ .

Generally speaking, either the full multinomial regression or conditional regression, though describing a much more detailed PMD pattern, could suffer from an overfitting issue when the data is limited, while the simpler regression model in the main text shows an accept-

690

able statistic power even with extremely small amount of data [A figure to cite?], we thus  
recommend the readers to use the simpler regression model when less data is applied.

692

## D | PMDTOOLS

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We use a way introduced by (Skoglund et al., 2014) to fish out the ancient strands with intensive PMD patterns from samples.

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702

According to (Skoglund et al., 2014), three nonmutually exclusive events can lead to an observation of  $C \rightarrow T$  or  $G \rightarrow A$ , namely (i) a true biological polymorphism (occurring at rate  $\pi$ ), (ii) a sequence error (rate  $\epsilon$ , can be extracted from the base quality scores of the site on the sampled strand), and (iii) in the case of damaged DNA, the damaged nucleotide frequencies are assumed to be only related to its position from either termini of the ancient fragment ( $C \rightarrow T$  from 5' end, and  $G \rightarrow A$  from 3' end),

704

$$D_x = C + p(1 - p)^{|x|}, \quad (20)$$

706

where  $C = 0.01$  and  $p = 0.3$  are both constants.

708

710

The observation "Match" is defined as the case when we observe a  $C$  at a position whose reference is also a  $C$  or a  $G$  at a position whose reference is also a  $G$ . And the observation "Mismatch" represents the situation when we get a  $T$  or an  $A$  at a position whose reference nucleotide is a  $C$  or a  $G$ , respectively. The likelihoods of whether or not a specific fragment is damaged given the observation are calculated in the subsequent subsections.

### Model with PMD

If a strand is damaged, the probability that we observe a "Match" event at position  $x$  of this strand can be viewed as the sum of probabilities of three mutually exclusive events: (i) no biological difference between the reference and the sampled nucleotide, no damage and no sequencing error, (ii) no biological difference, damaged but the sequencing error lead to a "Match" observation, and (iii) no damage, and both the sequencing error and the biological

718 divergence contribute to a "Match" observation,

720 
$$P(\text{Match} | x, \text{PMD}) = (1 - \pi)(1 - \epsilon)(1 - D_x) + (1 - \pi)\epsilon D_x + \pi\epsilon(1 - D_x), \quad (21)$$

722 
$$P(\text{Mismatch} | x, \text{PMD}) = 1 - P(\text{Match} | x, \text{PMD}). \quad (22)$$

724 The likelihood that the focal strand is damaged given the observation at position  $x$  is  $S_x$  can then be calculated as follows,

726 
$$L(\text{PMD} | S_x) = \chi_{S_x=\text{Match}} P(\text{Match} | x, \text{PMD}) + \chi_{S_x=\text{Mismatch}} P(\text{Mismatch} | x, \text{PMD}), \quad (23)$$

728 where  $\chi_{S_x}$  is an indicator function.

730 **Model without PMD**

732 Similarly, the "Match" event at position  $x$  in the case without PMD (the NULL model) can be decomposed as two exclusive events: (i) no biological divergence and no sequencing error, or (ii) both biological divergence and sequencing error contribute to a "Match" observation. And we have the following equations,

734 
$$P(\text{Match} | x, \text{NULL}) = (1 - \pi)(1 - \epsilon) + \pi\epsilon \quad (24)$$

736 
$$P(\text{Mismatch} | x, \text{NULL}) = 1 - P(\text{Match} | x, \text{NULL}) \quad (25)$$

738 
$$L(\text{NULL} | S_x) = \chi_{S_x=\text{Match}} P(\text{Match} | x, \text{NULL}) + \chi_{S_x=\text{Mismatch}} P(\text{Mismatch} | x, \text{NULL}) \quad (26)$$

740 Skoglund et al., 2014 defines the likelihood ratio of a strand between the PMD model and the NULL model as its postmortem damage score (PMDS),

$$\text{PMDS} = \log \left\{ \frac{\prod_x L(\text{PMD} | S_x)}{\prod_x L(\text{NULL} | S_x)} \right\}, \quad (27)$$

The strands with the PMDS exceeding a empirical p-value threshold (???) will be fished out as intensively damaged fragments.



### **3** *Paper II*

The following pages contain the paper:

**Christian Michelsen**, Christoffer C. Jørgensen, Mathias Heltberg, Mogens H. Jensen, Alessandra Lucchetti, Pelle B. Petersen, Troels C. Petersen, Henrik Kehlet (2022). “Preoperative prediction of medical morbidity after fast-track hip and knee arthroplasty – a machine learning based approach”.

Manuscript (Title Page, Abstract, Body, References, Figure  
Legends)

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## Preoperative prediction of medical morbidity after fast-track hip and knee arthroplasty - a machine learning based approach.

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4. Clinical Trial Number: The Centre for Fast-track Hip and Knee Replacement Database was registered  
as a study registry on ClinicalTrials.gov:NCT01515670

5. Prior presentations: Not applicable

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4     **6. Acknowledgements:** The members of the Centre for Fast-track Hip and Knee Replacement Database  
5     collaborative group all contributed by implementing the fast-track protocol at their respective departments  
6     and reviewing the final manuscript.  
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22     **7. Word and Element Counts:**  
23    Abstract: 300/300 Introduction: 466/500 Discussion:1278/1500 Figures:3 Tables:2 Appendices:2  
24    Supplementary Digital Files:4  
25     **8. Abbreviated title:** Machine learning models in joint replacement  
26     **9. Summary Statement:** Not applicable.  
27     **10. Funding:** The study received funding from the Lundbeck Foundation, Denmark, as well as from  
28     institutional and departmental sources.  
29     **11. Conflict of interest:** Prof. Kehlet is a board member of "Rapid Recovery", by Zimmer Biomet. Mr.  
30     Heltberg is sponsored by a grant from the Lundbeck Foundation, independently of the present study.  
31     Dr. Petersen is an advisory member of Sanofi outside of the present study.  
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## Abstract

**Background:** Introduction of machine-learning models has potentially improved prediction of postoperative hospitalization and morbidity after hip and knee replacement. However, few studies include enhanced recovery programs, and most rely on administrative coding with limited follow-up and information on perioperative care. Thus, benefits of machine-learning models for prediction of postoperative morbidity in enhanced recovery hip and knee replacement remain uncertain.

**Methods:** Multicenter cohort study from 2014-2017 in enhanced recovery total hip and knee replacement. Prospective recording of comorbidity and prescriptions. Information on length of stay and readmissions through the Danish National Patient Registry and medical records. Data was split into training (n:18013) and test sets (n:3913). A machine-learning model with 33 variables was used for predicting “medical” morbidity with a length of stay of >4 days or 90-days readmission and compared to a full logistic regression model. In addition, a machine-learning model excluding age, an age-only model and parsimonious machine-learning and logistic regression models using the ten most important variables were evaluated. Model performances were evaluated using several metrics, including precision, operating receiver (AUC) and precision recall curves (AUPRC). Variable importance was analyzed using Shapley Additive Explanations values.

**Results:** With 782 (20%) “risk-patients”, precision, AUC and AUPRC were 13.6%, 76.3% and 15.5% for the full and 12.8%, 75.9% and 17.1% for the parsimonious machine-learning models vs. 12.5%, 74.5% and 15.7% for the full logistic regression model. The machine-learning model excluding age and the Age-only model performed worse. Of the top ten variables, eight were shared between the full machine-learning and logistic regression models, and the importance of specific prescribed drugs varied considerably with age.

**Conclusion:** A machine-learning algorithm using preoperative characteristics and prescriptions likely improves identification of patients in high-risk of medical complications after fast-track hip and knee replacement. Such algorithms could help identify patients who benefit from intensified perioperative care.

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## INTRODUCTION

Prediction of postoperative morbidity and requirement for hospitalization is important for planning of health care resources. With regard to the common surgical procedures of primary total hip and knee arthroplasty, the introduction of enhanced recovery or fast-track programs has led to a significant reduction of postoperative length of stay (length of stay) as well as morbidity and mortality.<sup>1-3</sup> However, despite such progress, a fraction of patients still have postoperative complications leading to prolonged length of stay or readmissions.<sup>1,3,4</sup> Consequently, in order to prioritize perioperative care, many efforts have been published to preoperatively predict length of stay and morbidity using traditional risk factors such as age, preoperative cardio-pulmonary disease, anemia, diabetes, frailty, etc.<sup>4-8</sup> These efforts have been based on traditional statistical methods, most often multiple regression analyses, and essentially concluding that it is “better to be young and healthy than old and sick”. Consequently, despite being statistically significant, conventional risk-stratification based on such studies has had a relatively limited clinically relevant ability to predict and reduce potentially preventable morbidity and length of stay.<sup>4-8</sup> More recently, machine-learning methods have been introduced with success in several areas of healthcare and where preliminary data suggest them to improve surgical risk prediction compared to traditional risk calculation in certain anesthetic and surgical conditions.<sup>9,10</sup> This is also the case in total hip replacement, total knee replacement and uni-compartmental knee replacement, where several publications on machine-learning algorithms for prediction of length of stay,<sup>11,12</sup> complications,<sup>13</sup> disability,<sup>14</sup> potential outpatient setup,<sup>15</sup> readmissions<sup>16</sup> or payment models,<sup>17,18</sup> have shown promising predictive value compared to conventional statistical methods.<sup>19</sup> However, few papers have included enhanced recovery programs, and most are based on large database cohorts with the presence of risk factors and complications often relying on administrative coding with limited information on perioperative care, follow-up and discharge destination. In our previous study of 9512 total hip and knee replacements within an enhanced recovery protocol and including the above information, we did not find advantages of machine-learning methods compared to logistic regression in predicting a length of stay > 2 days.<sup>20</sup> However, this may have been due to data imbalance, lack of details on medication and the chosen outcome of length of stay of >2 days.<sup>20</sup> Thus, machine-learning models remain promising and could provide an improved basis for identifying a potential “high-risk” surgical

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4 population who may benefit from more extensive preoperative evaluation and postoperative  
5 medical care.  
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7 Consequently, we analyzed whether an improved machine-learning model was better for  
8 preoperative prediction of medical complications resulting in prolonged length of stay and  
9 readmissions compared to a traditional logistic regression model, in a large consecutive cohort  
10 of patients undergoing fast-track total hip and knee replacement within a national public health-  
11 care system.<sup>1</sup> In addition to well-defined patient-reported preoperative risk-factors, we also  
12 included information on dispensed reimbursed prescriptions 6 months prior to surgery using a  
13 nationwide registry.<sup>21</sup>  
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## 21 Method

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25 Reporting of the study is done in accordance with the Transparent reporting of multivariable  
26 prediction model for individual prognosis or diagnosis (TRIPOD) statement<sup>22</sup> and the Clinical AI  
27 Research (CAIR) checklist proposal.<sup>23</sup>  
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29 The study is based on the Centre for Fast-track Hip and Knee Replacement database which is a  
30 prospective database on preoperative patient characteristics and enrolling consecutive patients  
31 from 7 departments between 2010 and 2017. The database is registered on ClinicalTrials.gov  
32 as a study registry (NCT01515670). Permission to review and store information from medical  
33 records without informed consent was acquired from Center for Regional Development (R-  
34 20073405) and the Danish Data Protection Agency (RH-2007-30-0623). Patients completed a  
35 preoperative questionnaire with nurse assistance if needed. Additional information on  
36 reimbursed prescriptions 6 months prior to surgery was acquired using the Danish National  
37 Database of Reimbursed Prescriptions (DNDRP) which records all dispensed prescriptions with  
38 reimbursement in Denmark.<sup>21</sup> Finally, data were combined with the Danish National Patient  
39 Registry (DNPR) for information on length of stay (counted as postoperative nights spent in  
40 hospital), 90-days readmissions with overnight stay and mortality. In case of length of stay >4  
41 days or readmission, patient discharge summaries were reviewed for information on  
42 postoperative morbidity and in case of insufficient information, the entire medical records were  
43 reviewed. Readmissions were only included if considered related to the surgical procedure, thus  
44 excluding planned procedures like cancer workouts, cataract surgery, etc. Readmissions due to  
45 urinary tract infection or dizziness after day 30 were also considered unrelated to the surgical  
46 procedure. In case of postoperative mortality the entire medical record, including potential  
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4 readmissions, was reviewed to identify cause of death. Evaluation of discharge and medical  
5 records was performed by PP supervised by CJ. In case of disagreement, records were  
6 conferred with HK. Subsequently, causes of length of stay >4, readmissions or mortality were  
7 classified as “medical” when related to perioperative care (renal failure, falls, pain, thrombosis,  
8 anemia, venous thromboembolism or infection etc.) and “surgical” if related to surgical  
9 technique (prosthetic infection, revision surgery, periprosthetic fracture, hip dislocation, etc.).<sup>1</sup> In  
10 case of a length of stay 4-6 days with a standard discharge summary describing a successful  
11 postoperative course, it was assumed that no clinically relevant postoperative complications had  
12 occurred. If length of stay was >6 days but with standard discharge summary, the entire medical  
13 record was evaluated to confirm that no relevant complications had occurred.  
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15 For the present study, only cases between 2014 and 2017 were used to provide the most up-to  
16 date data. All patients had elective unilateral total hip and knee replacement in dedicated  
17 arthroplasty departments with similar fast-track protocols, including multimodal opioid sparing  
18 analgesia with high-dose (125mg) methylprednisolone, preference for spinal anesthesia, only in-  
19 hospital thromboprophylaxis when length of stay ≤5 days, early mobilization, functional  
20 discharge criteria and discharge to own home.<sup>1</sup> There was no selection criteria for the fast-track  
21 protocol as it is considered standard of care, but we excluded patients with previous major hip  
22 or knee surgery within 90-days of their total hip or total knee replacement and total hip  
23 replacement due to severe congenital joint disorder or cancer.  
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## Outcomes

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40 The primary outcome was to compare prediction quality when using a machine-learning model  
41 to predict the occurrence of “medical” complications resulting in a length of stay >4 days or  
42 readmission compared to a traditional logistic regression model (outcome A). Secondarily, we  
43 investigated how inclusion of cases with a length of stay >4 days but no reported “medical”  
44 complication as a positive outcome influenced the model (outcome B). For both outcomes, we  
45 also investigated whether a parsimonious model including only the top ten variables would  
46 perform equally well as the full model, and whether the effect of age per se would compare to  
47 the full machine-learning model. All figures and tables in the main text and Appendix are based  
48 on outcome A; the corresponding figures for outcome B are reported in the Supplemental Digital  
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## 4 Statistical Analysis 5

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Data was initially trimmed by removing 156 patients (1.7%) who were outliers with regards to weight (<30 kg or >250 kg) and height (<100 cm or >210 cm) or where these data were missing. To reduce the risk of overfitting, the dataset was subsequently split into a training set consisting of 18.013 (82.2%) procedures from 2014-2016 and a test set of 3913 (17.8%) procedures from 2017.

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As a reference model, we used classical logistic regression using all 33 input variables (table 1). Cases of missing values in the logistic regression model were handled by imputing missing values with the median of present values. All variables were then normalized.

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In addition, we used Boosted Decision Trees (LightGBM)<sup>24</sup> for the machine-learning models, as such methods work well with categorical data and missing values. We tried using both normal cross entropy and FocalLoss<sup>25</sup> as the objective function for the machine-learning model. The reason for testing FocalLoss was to allow the machine-learning model to focus more on the (few) positives.

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The full machine-learning model was trained and hyperparameter optimized using state of the art machine-learning methods. The models were trained on the training data and then used for making predictions on the unseen test data (see supplementary for details). The classification threshold was calibrated such that no more than 20% of the total number of patients were predicted as positive by the model (a positive predictive fraction (PPF) of 20%). We also included results for PPF values of 25% and 30%. Furthermore, we trained two parsimonious models using machine-learning and logistic regression with only the 10 most important features. Finally, we specifically explored the influence of increasing age, by constructing a model based only on age (Age), and a machine-learning model based on all variables except for age.

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To investigate the importance of the included variables, we computed the SHapley Additive exPlanations (SHAP) values, which provide estimates on which variables contribute most to the risk score predictions.<sup>26,27</sup> Finally, we investigated a potential relation between reimbursed prescribed cardiac drugs, anticoagulants, psychotropics and pulmonary drugs and age, as the relation between polypharmacy and postoperative outcomes have mainly been found in older patients.<sup>28</sup>

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For evaluating model performance, we computed the number of true positives, false positives, false negatives, true negatives, sensitivity (true positive rate), precision (positive predictive value). Since the data was quite imbalanced (about a 1:20 positive:negative ratio) we also computed the Matthews Correlation Coefficient (MCC) which is independent of class

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 4      imbalance.<sup>29,30</sup> The MCC ranges between -1 (the 100% wrong classifier), 0 (the random  
 5      classifier), and +1 (the perfect classifier). Finally, we computed the area under the receiver  
 6      operating characteristic curve (AUC) and the area under the precision recall curve (AUPRC). To  
 7      evaluate the statistical difference between the classifiers, we applied a Bayesian metric  
 8      comparison P(sensitivity),<sup>31</sup> which is the probability that a model will perform better than the  
 9      machine-learning model relative to the sensitivity. Thus, for two equally performing models  
 10     P(sensitivity) is ≈ 50%.  
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## Results

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 19      Median age in the 3913 patients was 70 years (IQR 62-76), 59% were female and 58% had  
 20      total hip replacement (table 1). Details on prescribed drug types are shown in Appendix 1.  
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 22      Median length of stay was 2 (IQR: 1-2) days with 7.6% 90-days readmissions and outcome A  
 23      occurring in 182 (4.7%) patients. When applying any model with a positive prediction fraction of  
 24      20% to the 3913 patients, 782 qualified as “risk-patients”. The results are summarized in figure  
 25      1 and table 2. When considering risk scores from the full machine-learning (figure 1a) and full  
 26      logistic regression model leading to this risk-patient selection, 106 and 98 had outcome A,  
 27      respectively. Correspondingly, the sensitivity and precision were 58.2% and 13.6% for the full  
 28      machine-learning and 53.8% and 12.5% for the full logistic regression model, respectively. The  
 29      full machine-learning model was superior (figure 1b) on essentially all parameters compared to  
 30      any of the other models, although the differences were minor (table 2). The results were similar  
 31      when using positive prediction fractions of 25% and 30%, but with the sensitivity for the full  
 32      machine-learning model increasing to 64.4% and 69.2% and precision decreasing to 12.0% and  
 33      10.7%, respectively (Appendix table 2).

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 35      Both the machine-learning model excluding age and age-only model had significantly lower  
 36      sensitivity than the full machine-learning model (figure 1b). Despite age being the single most  
 37      important variable (figure 2), the machine-learning model excluding age performed as well as  
 38      the age-only model.

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 40      When evaluating feature importance, we found a strong correlation between the full machine-  
 41      learning and full logistic regression model, with age and use of walking aids being the most  
 42      important variables in both (figure 2a). From the combined importance of variables outside the  
 43      top ten, the machine-learning approach extracted more information with fewer variables than  
 44      logistic regression (figure 1b).

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4 For the full machine-learning model, there was a clear signal that increasing age, number of  
5 reimbursed prescriptions, and presence of comorbidity, all contributed to an increased risk  
6 score. In contrast, a recent date of surgery and an increased hemoglobin level seemed to  
7 reduce the calculated risk (figure 2b). Individual analysis of the SHAP interaction values for  
8 types of anticoagulant prescriptions revealed that prescriptions on vitamin-K antagonists (VKA)  
9 or adenosine diphosphate (ADP) antagonists increased, while acetylic salicylic acid and direct  
10 oral anticoagulants (DOAC) reduced the risk score of the full machine-learning model,  
11 regardless of age (figure 3a). The SHAP analysis of prescribed cardiac drugs revealed that  
12 prescriptions on  $\text{Ca}^{2+}$ -antagonists and betablockers in combination with one or two other  
13 antihypertensives increased the risk-score, as did prescriptions on nitrates, other  
14 antihypertensives and antiarrhythmics. For the remaining cardiac drugs, prescriptions either  
15 reduced or had minor influence, and with limited relation with age (figure 3b). Preoperative  
16 psychotropic prescriptions increased the risk-score except for antipsychotics (0.6%). For users  
17 of selective serotonin inhibitors there was a clear age-related distinction with the risk score  
18 being increased in elderly patients but decreased in those < 60 years (figure 3c). Finally, the risk  
19 score increased with prescriptions on inhalation steroid and  $\beta$ -blockers, and more accentuated  
20 in the younger patients (figure 3d).

21 The results including patients with a length of stay >4 days, but no reported postoperative  
22 complications (outcome B) were similar as for outcome A. In general, we found that the full  
23 machine-learning model was superior to the others, although the difference were smaller than  
24 for outcome A. (Supplemental Digital Content table S1 listing outcome parameters and  
25 Supplemental Digital Content 2 figure S1a-b showing distributions and ROC curves for outcome  
26 B). While the ten most important variables for the full machine-learning model remained  
27 unchanged, familiar disposition for venous thromboembolism replaced gender as one of the top  
28 ten important variables in the full logistic regression model (Supplemental Digital Content figure  
29 S2a-b showing SHAP values for outcome B). Furthermore, the SHAP analysis on specific  
30 prescribed drugs demonstrated that the machine-learning model found no benefits from  
31 information on prescriptions on respiratory drugs, why all SHAP values were zero. In addition,  
32 the reduced risk with acetylsalicylic acid and DOAC prescriptions, as well as the influence of  
33 practically all cardiac drugs except for nitrates, other antihypertensives and antiarrhythmics, was  
34 attenuated (Supplemental Digital Content 4 figure S3a-d showing SHAP-values of prescriptions  
35 of specific drugs for outcome B).

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## Discussion

We found that using a machine-learning algorithm including all 33 available variables and a parsimonious machine-learning-algorithm encompassing only the 10 most important predictors improved prediction of patients at increased risk of having a length of stay >4 days or readmissions due to medical complications compared to traditional logistic regression models. In contrast, when also including patients having a length of stay >4 days but without a well-defined complication as an outcome, the parsimonious machine-learning model was slightly worse than a traditional logistic regression model including all variables. We also found that although age was the single most important predictor of both outcome A and B, it was less suited for prediction of postoperative medical complications after fast-track total hip and knee replacement on its own. Finally, we demonstrated how the chosen classification threshold of the machine-learning algorithm influenced model performance through an increase in sensitivity at the cost of decreased precision.

A previous systematic review also found that machine-learning algorithms may provide better prediction of postoperative outcomes in THA and TKA.<sup>32</sup> However, the authors concluded that such models performed best at predicting postoperative complications, pain and patient reported outcomes and were less accurate at predicting readmissions and reoperations.<sup>32</sup> That machine-learning algorithms may improve prediction of complications after THA and TKA compared to traditional logistic regression was also found by Shah *et al.* who used an automated machine-learning framework to predict selected major complications after THA.<sup>13</sup> However, theirs was a retrospective study based on diagnostic and administrative coding and the selected complications occurred only in 0.61% of patients, potentially limiting clinical relevance. In contrast, we aimed at identifying a cohort which would comprise 20% of patients in which we found about 60% of all medical complications. This we believe, is within the means of the Danish socialized healthcare system to allocate additional resources for intensified perioperative care and with both patient-related and economic benefits due to potentially avoided complications and costs.

In contrast to many other machine-learning studies,<sup>33</sup> our dataset included not only preoperative data but also only one paraclinical variable, which was preoperative hemoglobin. Although the inclusion of other laboratory tests such as preoperative albumin, sodium and alkaline phosphatase has been found to be of importance in machine-learning algorithms for home discharge in UKA<sup>12</sup> and spine surgery,<sup>9</sup> they are not standard in our fast-track protocols and not easy to interpret from a pathophysiological point of view. As there is a need to prioritize the

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4 limited health-care resources, most decisions on which patients may benefit from more  
5 extensive postoperative care will likely need to be conducted preoperatively. Thus, although  
6 postoperative information such as duration of surgery, perioperative blood length of stays or  
7 postoperative hemoglobin have been included in other studies<sup>33</sup>, we decided against the use of  
8 peri- and postoperative data. The same approach has been used by Ramkumar *et al.* who used  
9 U.S. National Inpatient Sample data including 15 preoperative variables, to predict length of  
10 stay, patient charges and disposition after both TKA<sup>34</sup> and THA.<sup>18</sup> However, these studies were  
11 not conducted in a socialized health care system, and the main focus was on the need for  
12 differentiated payment bundles and without specific information on the reason for increased  
13 length of stay or non-home discharge.<sup>34</sup> Wei *et al.* used an artificial neural network model to  
14 predict same-day discharge after TKA, based on the NSQUIP database from 2018 and found  
15 that six of the ten most important variables were the same compared with logistic regression,  
16 similar to our findings.<sup>35</sup> However, patients with one-day length of stay were intentionally  
17 excluded due to variations in in-patient vs. out-patient registration.<sup>35</sup>  
18  
19 Age has traditionally been a major factor when predicting surgical outcomes which is why we  
20 choose to specifically evaluate its effect on our risk-prediction. That age is important for risk-  
21 prediction was further illustrated by the machine-learning model without age being comparable  
22 to the age-only model. Note that, although elderly patients had increased risk of postoperative  
23 complications, likely related to decline of physical reserves,<sup>36</sup> the use of chronological age alone  
24 as a selection criteria for being a “risk-patient” was inferior compared to both machine-learning  
25 and logistic regression models incorporating comorbidity and functional status.  
26  
27 We used the SHAP values for estimation of feature importance, thus providing a better  
28 understanding of the otherwise “black-box” machine-learning model. The SHAP values showed  
29 which variables contribute most to the risk-score predictions.  
30  
31 Our inclusion of specific data on reimbursed prescriptions 6 months prior to surgery based upon  
32 the unique Danish registries, unsurprisingly found increased risk-scores with increased number  
33 of prescriptions and with the majority being in elderly patients. Similarly, a Canadian study in  
34 elective non-cardiac surgery found decreased survival and increased length of stay and  
35 readmissions and costs in patients >65 years with polypharmacy.<sup>28</sup> However, this is a complex  
36 relationship where some patients benefit from their treatments, while others may suffer from  
37 undesirable side-effects. Consequently, the authors cautioned against altering perioperative  
38 practices based on current evidence.<sup>28</sup> However, the information from the included prescriptions  
39 with SHAP analysis may provide inspiration for new hypothesis-generating studies such as  
40 investigation of the potential differences in risk-profile between having preoperative prescribed  
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4 VKA and DOAKs. Also, the age-related differences in risk from SSRI's seen in our study could  
5 guide further studies on "deprescription".  
6  
7 Another requirement for machine-learning-algorithms to be clinically useful is user friendliness  
8 and not depending on excessive additional data collection by the attending clinicians. In this  
9 context, it was a bit disappointing that the parsimonious machine-learning algorithm with only  
10 the ten most important variables was slightly worse at predicting outcome B than the full logistic  
11 regression model. A reason for this could be that when including a length of stay >4 days but  
12 without described medical complications, the combination of all variables provide information  
13 not available by merely including the ten most important ones. This highlights the need for as  
14 much detailed, and preferably non-binary, data as possible to fulfill the true potential of  
15 machine-learning algorithms.  
16  
17 Our study has some limitations. First, one of the strengths of machine learning compared to  
18 logistic regression is the analysis of multilevel continuous data, whereas we included only a  
19 limited number of, often binary, preoperative variables. This could have limited the full  
20 realization of our machine learning model. As previously mentioned, we excluded intraoperative  
21 information, including type of anesthesia, surgical approach etc. all of which may influence  
22 postoperative outcomes. The observational design of this study means that we cannot exclude  
23 unmeasured confounding or confounding by indication. Also, despite that the DNDRP has a  
24 near complete registration of dispensed medicine in Denmark, some types or drugs, especially  
25 benzodiazepines, are exempt from general reimbursement and thus not sufficiently captured.<sup>21</sup>  
26 Furthermore, it is doubtful whether the patients used all types of drugs at the time of surgery  
27 (e.g. heparin which is rarely for long-term use). Finally, classification of a complication being  
28 "medical" depended on review of the discharge records which can also introduce bias. However,  
29 we believe our approach to be superior to depending only on diagnostic codes which often are  
30 inaccurate<sup>37</sup> and provide limited details on whether the complication may be attributed to a  
31 medical or surgical adverse event. The strengths of our study include the use of national  
32 registries with high degree of completion (>99% of all somatic admissions in case of the  
33 DNDRP),<sup>38</sup> prospective recording of comorbidity, extensive information on prescription patterns  
34 6 months prior to surgery and similar established enhanced recovery protocols in all  
35 departments.  
36  
37 In summary, our results suggest that machine-learning-algorithms likely provide clinically  
38 relevant improved predictions for defining patients in high-risk of medical complications after  
39 fast-track THA and TKA compared to a logistic regression model. Future studies could benefit  
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4 from using such algorithms to find a manageable population of patients who benefit from  
5 intensified perioperative care.  
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## Figure legends

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### Figure 1a-b

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- 1a) Distribution of full machine learning model risk scores for patients +/- outcome A. The dashed line marks the classification threshold of 20% positive prediction fraction.  
1b) Receiver operating curves (ROC) for the full machine learning model (F-MLM), full logistic regression model (F-LRM), parsimonious machine learning model (P-MLM), parsimonious logistic regression model (P-LRM), machine learning excluding age (MLM -age) and the age-only model (AM).

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### Figure 2a-b

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- 2a) The overall importance of the 10 most important variables measured by the SHAP-values for the full machine-learning and full logistic regression models on outcome A (LOS >4 days or readmission due to "medical" morbidity). Only the importance of prescribed anticoagulants and gender differ between the models. The contributions of the remaining variables are summed in the bottom bar.  
2b) The SHAP-values for the full machine-learning model on outcome A, where positive increase and negative values decrease the risk score. The color is related to the value of the variable with blue being lowest and red highest and each dot represents a patient.

### Figure 3a-d

3HAP scatter-plot on the contributions to the full machine-learning model on outcome A (LOS >4 days or readmission due to "medical" morbidity), for individual types of prescribed anticoagulants, cardiac drugs, psychotropics and respiratory drugs stratified by age.

#### 3a) Prescribed anticoagulants

VKA: vitamin K antagonists ASA: acetylsalicylic acid DOAC: direct oral anticoagulant ADP: Adenosine diphosphate ACE: angiotensin converting enzyme

#### 3b) Prescribed cardiac drugs

ACE: angiotensin converting enzyme AHT: antihypertensive. Other AHT were defined as AHT different from diuretics ANG-II/ACE inhibitors or Ca<sup>2+</sup>antagonists. IHD: Ischemic heart disease

#### 3c) Prescribed psychotropics

SSRI: Selective serotonin inhibitor SNRI: Serotonin and norepinephrine reuptake inhibitor NaRI: Norepinephrine reuptake inhibitor NaSSA: Norepinephrine and specific serotonergic antidepressants. AD: antidepressants BZ: Benzodiazepines (likely underreported due to limited general reimbursement in Denmark). ADHD: Attention-deficit/hyperactivity disorder

#### 3d) Prescribed respiratory drugs

SABA: Short-acting beta agonist LABA: long-acting beta agonist LAMA: Long-acting muscarinic antagonist.

table1

Table 1. patient demographics with and without outcome A (length of stay >4 days or readmissions due to "medical" morbidity) in the combined test and training dataset.

Preoperative characteristics n (%) unless otherwise specified	+outcome A (n:1180)	-outcome A (n:20837)
mean age (SD)	75.0 (68.0-81.0)	69.0 (62.0-75.0)
mean number of reimbursed prescriptions <sup>1</sup> (SD)	3.0 (1.0-4.0)	2.0 (0.0-3.0)
female gender	755 (64.0)	12133 (58.2)
Total hip replacement	636 (53.9)	11542 (55.4)
mean weight in kg (SD)	78.0 (67.0-91.0)	81 (70.0-93.0)
mean height in cm (SD)	168 (162.0-175.0)	170.0 (164.0-178.0)
mean body mass index (SD)	27.3 (23.9-31.2)	27.5 (24.6-31.1)
regular use of walking aid	552 (46.8)	4398 (21.5)
missing	29 (2.5)	359 (1.7)
living alone	578 (49.0)	6717 (32.2)
with others	571 (48.4)	13869 (66.6)
institution	24 (2.0)	113 (0.5)
missing	7 (0.6)	138 (0.7)
hemoglobin	8.2 (7.7-8.8)	8.6 (8.1-9.2)
missing	11 (0.9)	314 (1.5)
>2 units of alcohol/day	79 (6.7)	1589 (7.6)
missing	10 (0.8)	174 (0.8)
active smoker	130 (11.0)	2751 (13.2)
missing	11 (0.9)	141 (0.7)
cardiac disease	306 (25.9)	2750 (13.2)
missing	8 (0.8)	153 (0.7)
hypercholesterolemia	467 (39.6)	6062 (29.1)
missing	8 (0.7)	120 (0.6)
hypertension	738 (62.5)	10141 (48.7)
missing	64 (5.4)	663 (3.2)
pulmonary disease	182 (15.4)	1841 (8.8)
missing	5 (0.4)	96 (0.5)
previous cerebral attack	165 (14.0)	1086 (5.2)
missing	25 (2.1)	282 (1.4)
previous VTE	133 (11.3)	1481 (7.1)
missing	26 (2.2)	325 (1.6)
malignancy (undefined)	557 (47.2)	8843 (42.4)
previous radically treated malignancy	127 (10.8)	2065 (9.9)
missing	14 (1.2)	162 (0.8)
chronic kidney disease	50 (4.2)	273 (1.3)
missing	35 (3.0)	292 (1.4)
family member with VTE	155 (13.1)	2510 (12.0)
missing	1190 (16.1)	2569 (12.3)
regular snoring	266 (22.5)	5522 (26.5)
uncertain about snoring	208 (17.6)	3781 (18.1)
missing	259 (21.9)	3309 (15.9)
not feeling rested	468 (39.7)	9340 (44.8)

uncertain about being rested	48 (4.1)	809 (3.9)
missing	105 (8.9)	1230 (5.9)
psychiatric disorder	156 (13.2)	1590 (7.6)
missing	62 (5.3)	703 (3.4)
Characteristic based on combination of questionnaire and DNDRP		
<u>diabetes</u>		
diet treated diabetes <sup>2</sup>	29 (2.5)	274 (1.3)
oral antidiabetics	137 (11.6)	1448 (6.9)
insulin treated diabetes <sup>3</sup>	60 (5.1)	413 (2.0)
missing	7 (0.6)	98 (0.5)

SD: standard deviation VTE: venous thromboembolic event DNDRP: Danish National Database of Reimbursed Prescriptions.

<sup>1</sup>Antirheumatica, steroids, anticoagulants, cardiac, cholesterol lowering, respiratory and psychotropic drugs.

<sup>2</sup>Reported diabetes but no registered prescriptions <sup>3</sup>+/- oral antidiabetics

table2

Table 2: Performance of the six different models with a predefined positive prediction fraction of 20% for outcome A

Positive prediction fraction 20%	TP	FP	FN	TN	sensitivity	precision	MCC	AUROC	AUPRC	P (sensitivity)
Full machine-learning model	106	676	76	3055	58.2%	13.6%	21.1%	76.3%	15.5%	-
Full logistic regression model	98	684	84	3047	53.8%	12.5%	18.7%	74.5%	15.7%	19.7%
Parsimonious machine-learning model	100	682	82	3049	54.9%	12.8%	19.3%	75.9%	17.3%	26.1%
Parsimonious logistic regression model	95	687	87	3045	52.2%	12.1%	17.8%	73.7%	13.6%	12.4%
machine-learning model excluding age	88	694	94	3037	48.4%	11.3%	15.7%	72.3%	13.6%	3.1%
Age-only model	87	676	95	3055	47.8%	11.4%	15.8%	69.7%	12.1%	2.3%

TP: true positives FP: false positives FN: false negatives TN: true negatives MCC: Matthews correlation coefficient  
AUC: area under the ROC curve AUPRC: area under the precision recall curve P(sensitivity): probability that the model performs better than the machine-learning model relative to sensitivity.

Appendix1 (to appear in print publication)

## Appendix table 1

Details on specific drugs with reimbursed prescriptions 6 months preoperatively.  
Numbers are n (%)

		+Outcome A	-Outcome A
<b>Anticoagulants</b>			
none		679 (57.5)	15844 (76.0)
VKA		106 (9.0)	750 (3.6)
Heparin & Acetylsalicylic acid		0 (0.0)	7 (0.0)
DOAC		48 (4.1)	659 (3.2)
Acetylsalicylic acid		205 (17.4)	2492 (12.0)
Dipyradimol		5 (0.4)	29 (0.1)
ADP-antagonist		75 (6.4)	569 (2.7)
Acetylsalicylic acid & Dipyradimol		17 (1.4)	168 (0.8)
VKA & Acetylsalicylic acid		10 (0.8)	78 (0.4)
DOAC & Acetylsalicylic acid		6 (0.5)	41 (0.2)
VKA & ADP-antagonist		4 (0.3)	11 (0.1)
DOAC & ADP-antagonist		3 (0.3)	14 (0.1)
VKA & Heparin		1 (0.1)	21 (0.1)
DOAC & Acetylsalicylic acid & ADP-antagonist		1 (0.1)	3 (0.0)
Acetylsalicylic acid & ADP-antagonist		18 (1.5)	132 (0.6)
Acetylsalicylic acid & ADP-antagonist & Heparin		1 (0.1)	12 (0.1)
Acetylsalicylic acid & ADP-antagonist & Dipyradimol		1 (0.1)	7 (0.0)
<b>Cardiac prescriptions</b>			
none		321 (27.2)	9200 (44.2)
diuretics		77 (6.5)	1184 (5.7)
angiotensin-II/ACE-inhibitors		132 (11.2)	2683 (12.9)
Ca <sup>2+</sup> antagonists		55 (4.7)	773 (3.7)
β-blocker		29 (2.5)	559 (2.7)
nitrates		1 (0.1)	18 (0.1)
other antihypertensives		0 (0.0)	12 (0.1)
other types of medication for IHD		2 (0.2)	21 (0.1)
2 antihypertensives		177 (15.0)	2696 (12.9)
β-blocker & 1 antihypertensive <sup>1</sup>		92 (8.1)	1069 (5.1)
3 antihypertensives		50 (4.2)	548 (2.6)
β-blocker & 2 antihypertensives <sup>1</sup>		95 (8.1)	975 (4.7)
β-blocker & 3 antihypertensives <sup>1</sup>		25 (2.1)	265 (1.3)
4 antihypertensives		2 (0.2)	18 (0.1)
β-blocker & 4 antihypertensives		2 (0.2)	19 (0.1)
other antihypertensive & antihypertensives <sup>1</sup>		9 (0.8)	87 (0.4)
nitrates & any hypertensive		49 (4.2)	331 (1.6)
other drugs for IHD & any antihypertensive and/or nitrate		5 (0.4)	15 (0.1)
other antiarrhythmics & any antihypertensives		57 (4.8)	364 (1.7)
<b>Anticholesterols</b>			
none		708 (60.0)	14719 (70.6)
statins		457 (38.7)	5866 (28.2)
other anti-lipids		7 (0.6)	135 (0.6)
Statins +other anti-lipids		8 (0.7)	117 (0.6)

1	<b>Systemic steroids</b>	123 (10.4)	1149 (5.5)
2	<b>Antirheumatics</b>		
3	none	1143 (96.9)	20388 (97.8)
4	disease-modifying antirheumatic drugs	37 (3.1)	446 (2.1)
5	other antirheumatics	0 (0.0)	3 (0.0)
6	<b>Respiratory prescriptions</b>		
7	none	1000 (84.7)	18754 (90.0)
8	SABA	13 (1.1)	276 (1.3)
9	LABA or LAMA	19 (1.6)	217 (1.0)
10	inhalation steroid only	8 (0.7)	211 (1.0)
11	SABA & Ipratropium (+/- others)	6 (0.5)	18 (0.1)
12	LABA & steroid	45 (3.8)	474 (2.3)
13	LABA & LAMA & steroid	19 (1.6)	122 (0.6)
14	LAMA & steroid	0 (0.0)	11 (0.1)
15	LABA & LAMA	7 (0.6)	80 (0.4)
16	other pulmonary drugs	3 (0.3)	32 (0.2)
17	other pulmonary drugs & steroid	9 (0.8)	98 (0.5)
18	SABA & LABA or LAMA	6 (0.5)	96 (0.5)
19	SABA & LABA or LAMA & steroid	45 (3.8)	448 (2.2)
20	<b>Psychotropic prescriptions</b>		
21	none	952 (80.7)	18657 (89.5)
22	SSRI/SNRI/NaRI	100 (8.5)	1164 (5.6)
23	other antidepressants	1 (0.1)	17 (0.1)
24	antipsychotics	8 (0.7)	116 (0.6)
25	benzodiazepines <sup>2</sup>	0 (0.0)	7 (0.0)
26	anti-cholinergics or memantine	6 (0.5)	27 (0.1)
27	anti-ADHD drugs	1 (0.1)	10 (0.0)
28	NaSSA	25 (2.1)	184 (0.9)
29	other psychotropics	28 (2.4)	182 (0.9)
30	SSRI + other antidepressants	4 (0.3)	6 (0.0)
31	SSRI + NaSSA	8 (0.7)	94 (0.5)
32	SRRI + antipsychotics	11 (0.9)	87 (0.4)
33	SRRI + other psychotropics	7 (0.6)	84 (0.4)
34	benzodiazepines + any psychotropic	3 (0.3)	12 (0.1)
35	antipsychotics + any psychotropic	20 (1.7)	149 (0.7)
36	anti-ADHD + any psychotropic	0 (0.0)	14 (0.1)
37	NaSSA + any psychotropic	4 (0.3)	18 (0.1)
38	other psychotropics + any specified psychotropic	2 (0.2)	9 (0.0)

VKA: vitamin K antagonists DOAC: direct oral anticoagulant ADP: Adenosine diphosphate ACE: angiotensin converting enzyme IHD: Ischemic heart disease SABA: Short-acting beta agonist LABA: long-acting beta agonist LAMA: Long-acting muscarinic antagonist SSRI: Selective serotonin inhibitor SNRI: Serotonin and norepinephrine reuptake inhibitor NaRI: Norepinephrine reuptake inhibitor NaSSA: Norepinephrine and specific serotonergic antidepressants

<sup>1</sup>either diuretics, ACE/ANG-II inhibitors or Ca<sup>2+</sup>-antagonists <sup>2</sup>likely underreported due to limited general reimbursement for benzodiazepines in Denmark

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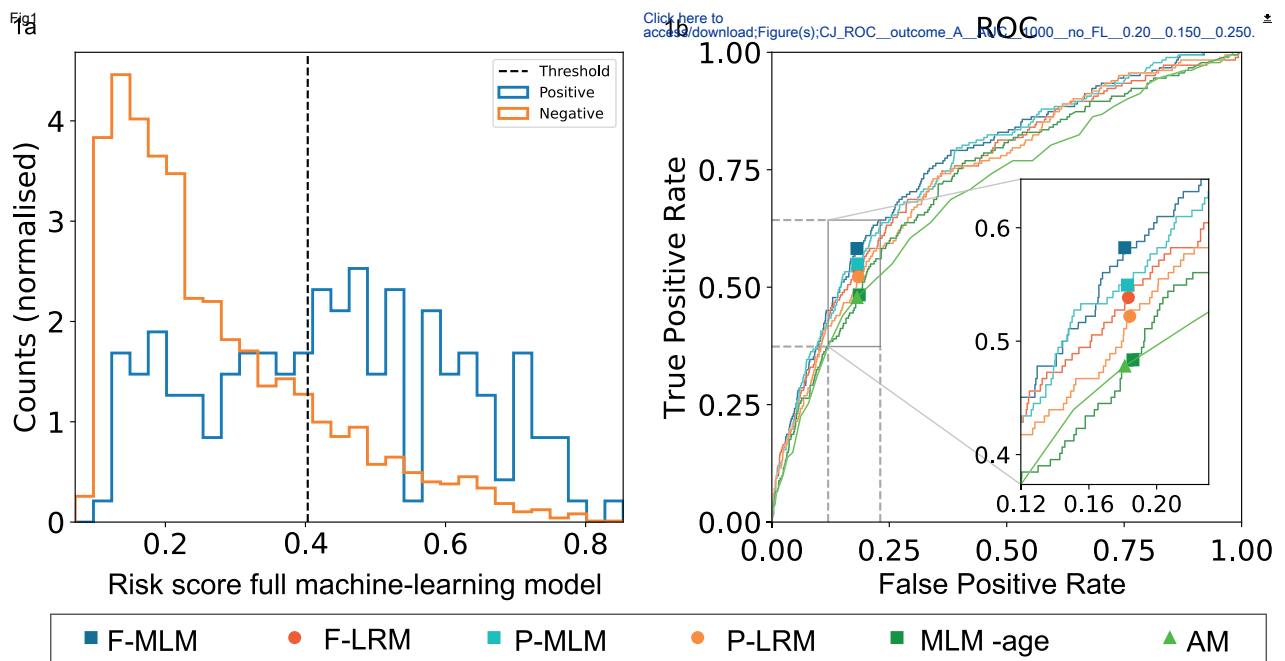
Appendix (to appear in print publication)

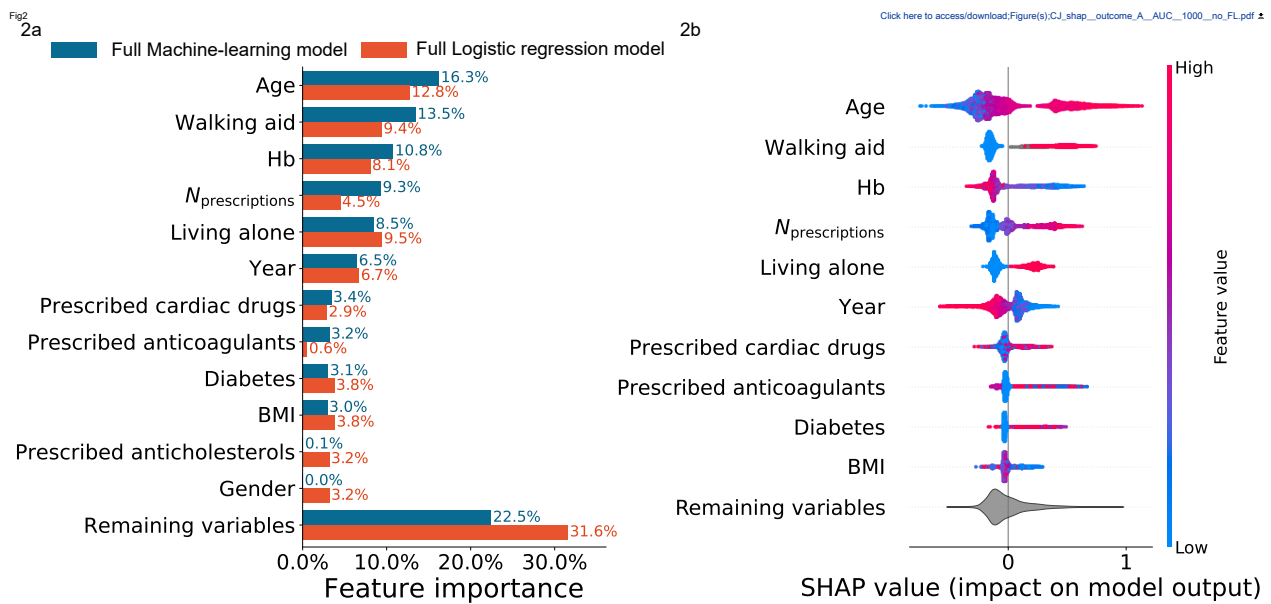
## Appendix table 2

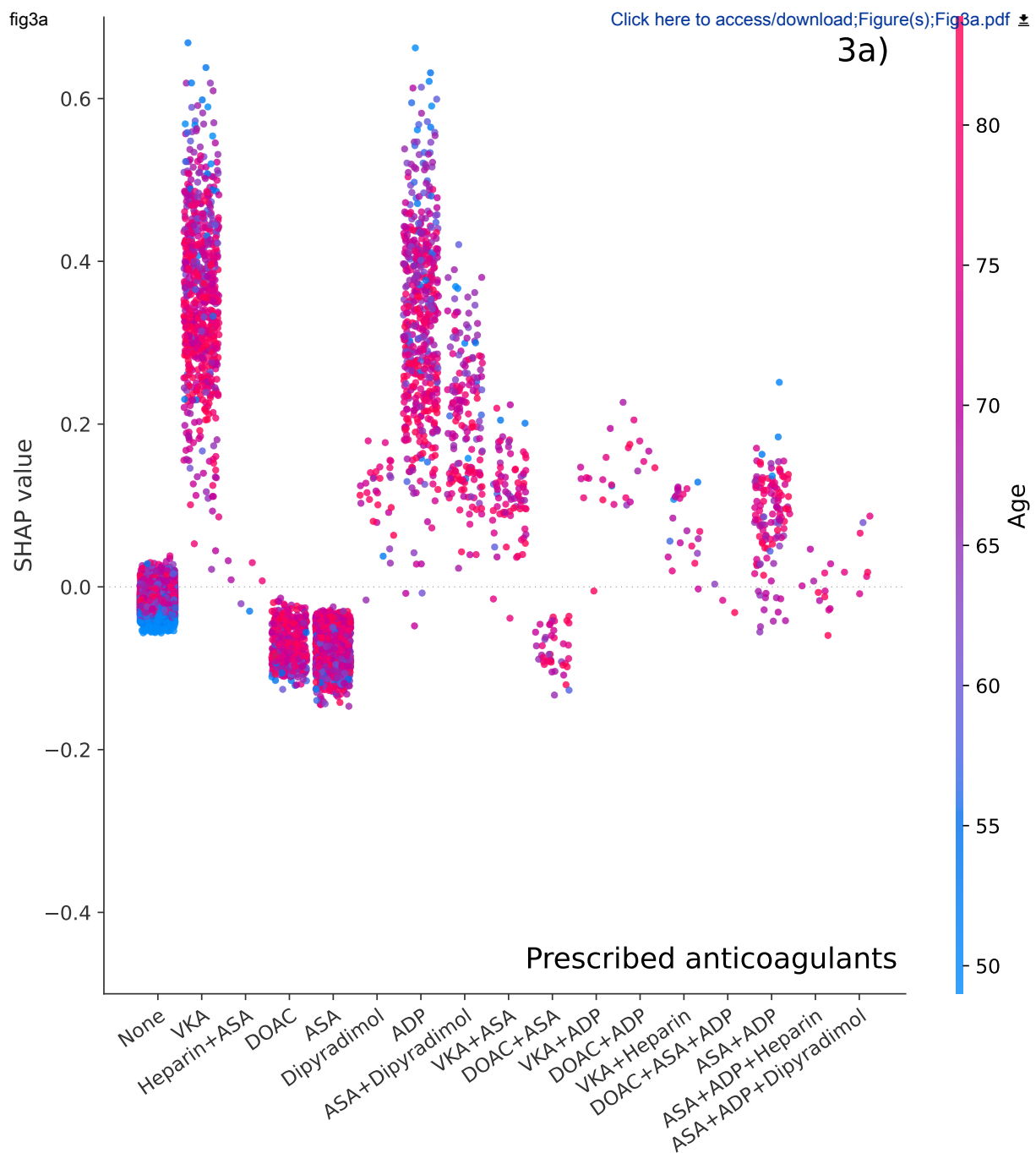
Performance of the six different models with a predefined positive prediction fraction of 25% and 30% for outcome A (LOS >4 days or readmission due to "medical" morbidity).

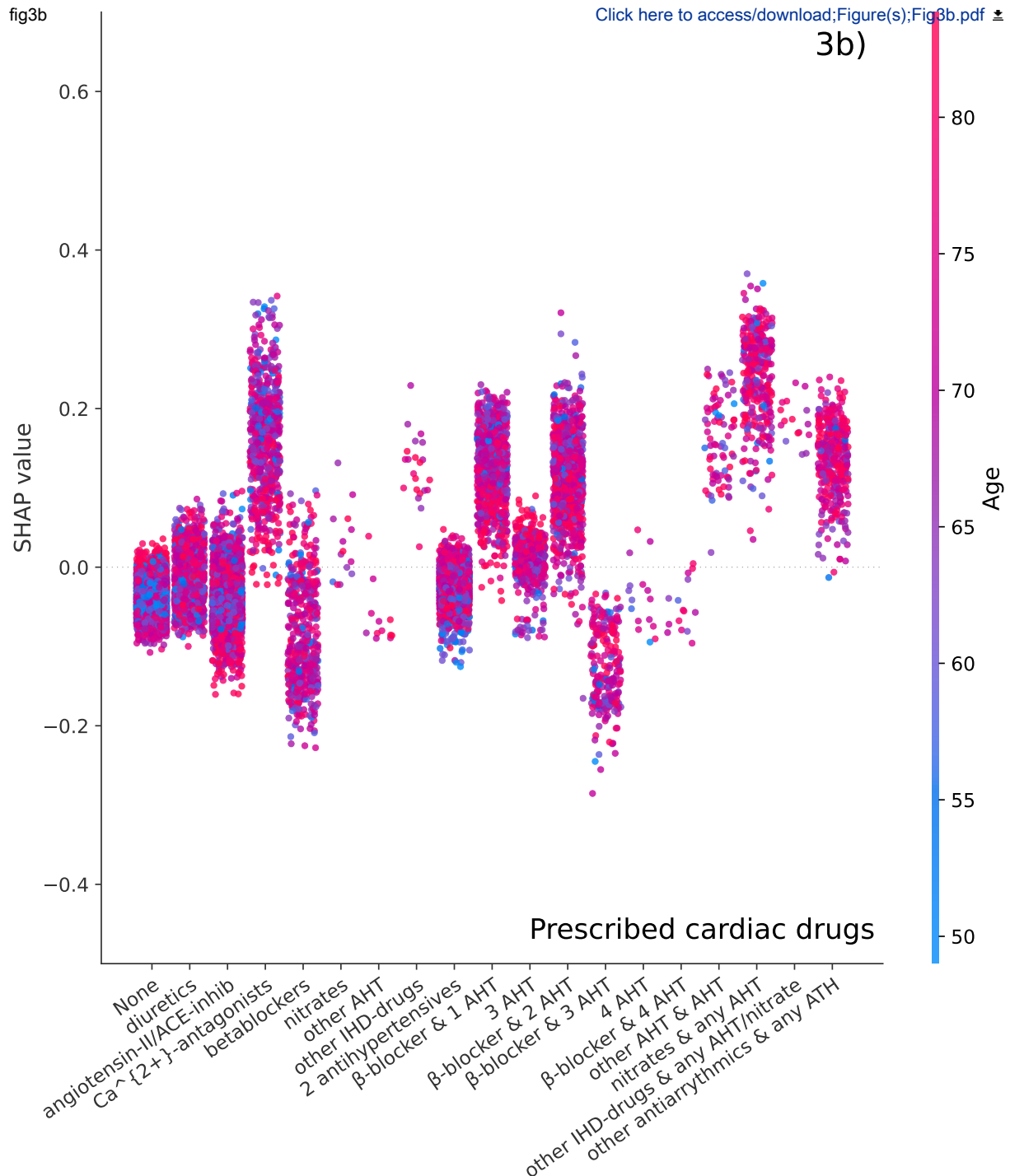
Positive prediction fraction 25%	TP	FP	FN	TN	sensitivity	precision	MCC	AUC	AUPRC	P (sensitivity)
Full machine-learning model	117	861	65	2870	64.3%	12.0%	20.0%	76.3%	15.5%	-
Full logistic regression model	110	868	72	2863	60.4%	11.2%	18.1%	74.5%	15.7%	23.1%
Parsimonious machine-learning model	115	863	67	2868	63.2%	11.8%	19.5%	75.9%	17.3%	41.2%
Parsimonious logistic regression model	106	872	76	2859	58.2%	10.8%	17.0%	73.4%	15.5%	11.8%
machine-learning model excluding age	106	872	76	2859	58.2%	10.8%	17.0%	72.3%	13.6%	11.8%
Age-model	94	824	88	2907	51.6%	10.2%	14.7%	69.7%	12.2%	0.7%
Positive prediction fraction 30%	TP	FP	FN	TN	sensitivity	precision	MCC	AUC	AUPRC	P (sensitivity)
Full machine-learning model	126	1047	56	2684	69.2%	10.7%	18.9%	76.3%	15.5%	-
Full logistic regression model	120	1053	62	2678	65.9%	10.2%	17.3%	74.5%	15.7%	25.2%
Parsimonious machine-learning model	124	1049	58	2682	68.1%	10.6%	18.4%	75.9%	17.3%	40.8%
Parsimonious logistic regression model	115	1058	67	2673	63.2%	9.8%	16.0%	73.7%	15.5%	11.1%
machine-learning model excluding age	116	1057	66	2674	63.7%	9.9%	16.3%	72.3%	13.6%	13.8%
Age-model	100	955	82	2776	54.9%	9.5%	13.9%	69.7%	12.2%	0.2%

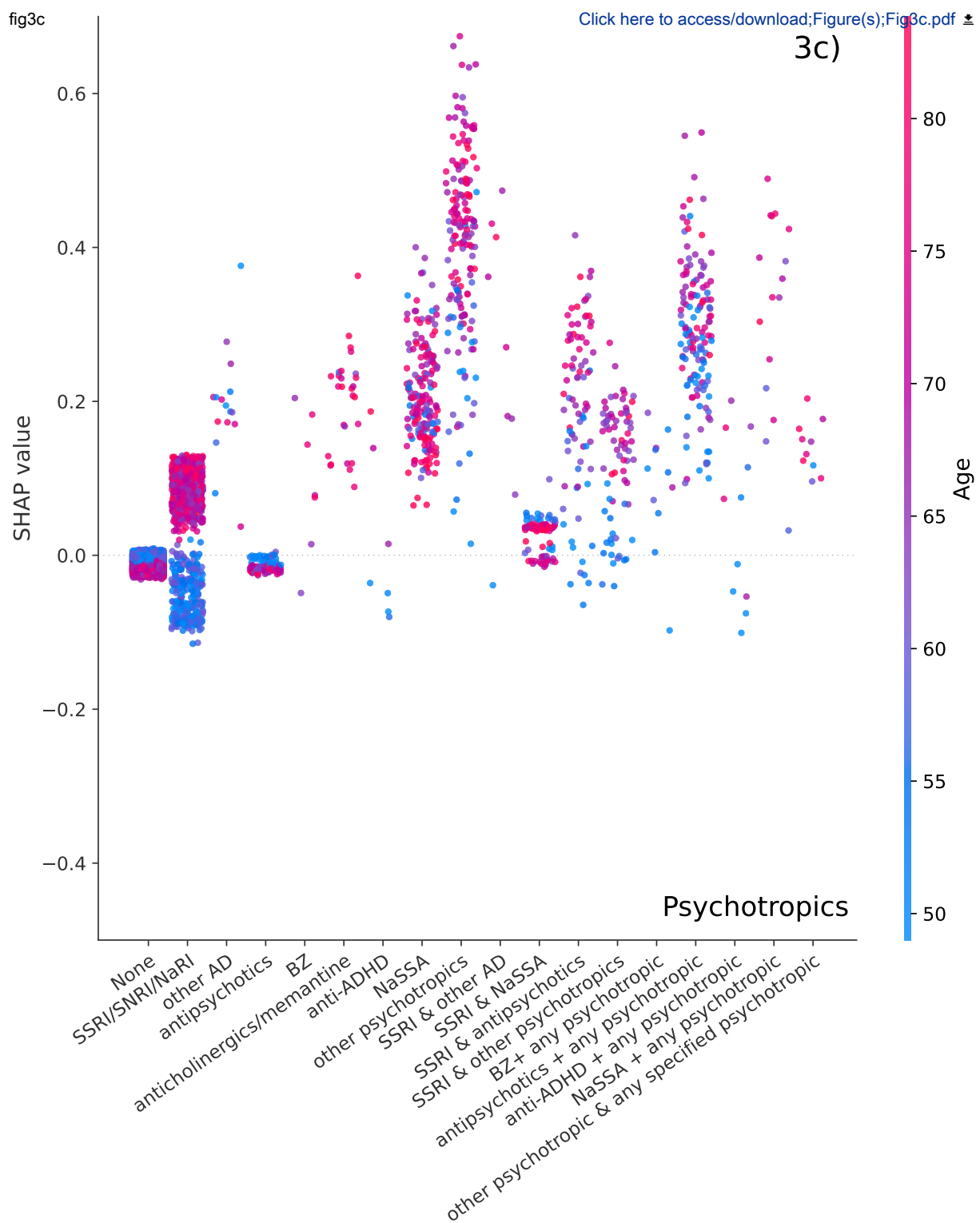
TP: true positives FP: false positives FN: false negatives TN: true negatives MCC: Matthews correlation coefficient AUC: area under the ROC curve AUPRC: area under the precision recall curve P(sensitivity): probability that the model performs better than the machine-learning model relative to sensitivity.

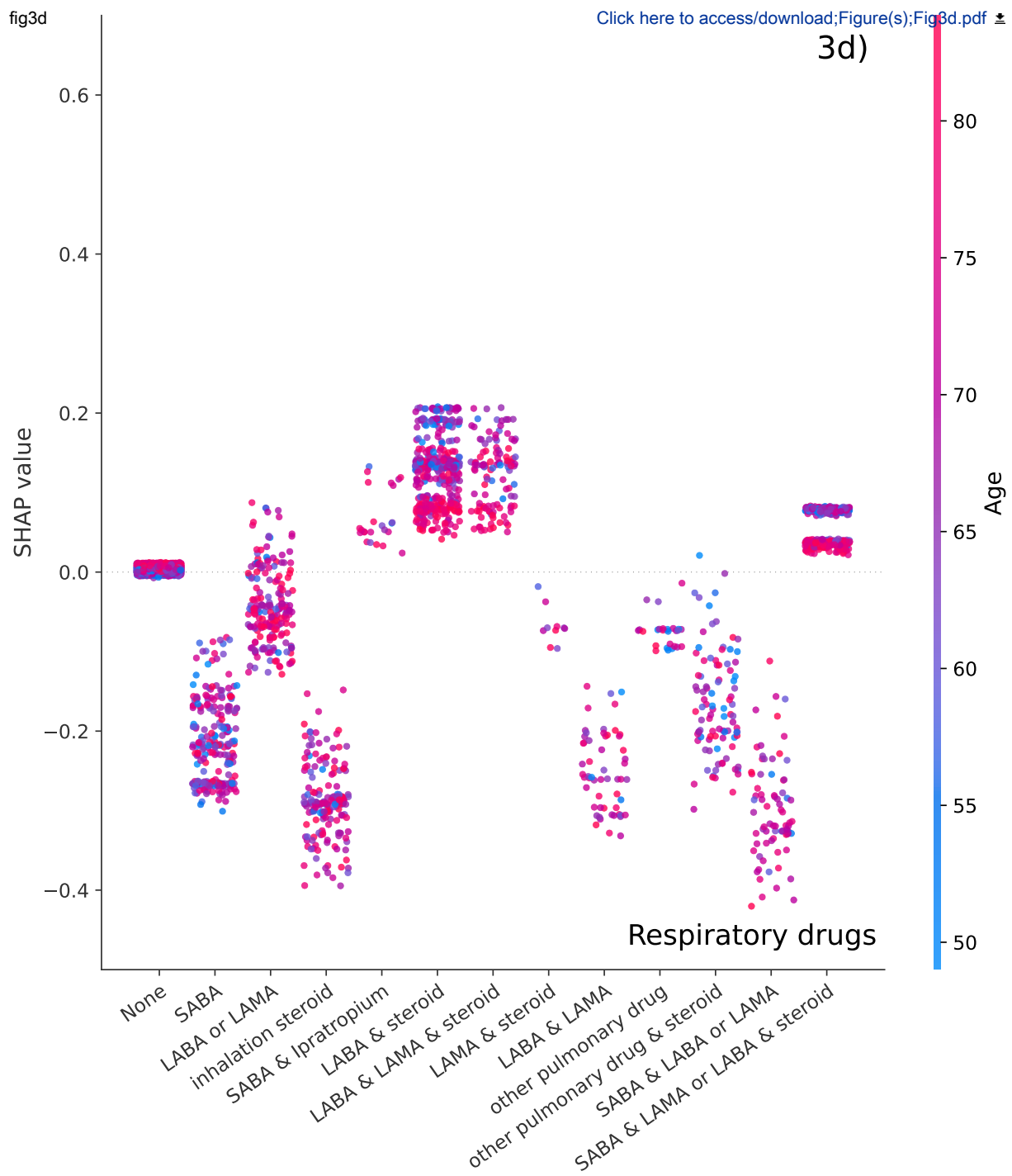












## Supplemental Digital Content 1

Table S1: Performance of different models for Outcome B (Los &gt;4 days or readmissions due to "medical" morbidity or LOS &gt;4 days but without recorded morbidity)

Positive prediction fraction 20%	TP	FP	FN	TN	sensitivity	precision	MCC	AUC	AUPRC	P (sensitivity)
Full machine-learning model	121	661	108	3023	52.8%	15.5%	20.5%	75.3%	17.1%	-
Full logistic regression model	115	667	114	3017	50.2%	14.7%	18.9%	74.1%	16.7%	28.3%
Parsimonious machine-learning model	111	671	118	3013	48.4%	14.2%	17.8%	74.4%	16.8%	17.2%
Parsimonious logistic regression model	109	673	120	3011	47.6%	13.9%	17.2%	73.1%	16.8%	12.9%
machine-learning model excluding age	110	672	119	3012	48.0%	14.1%	17.5%	72.8%	16.9%	15.1%
Age-model	102	661	127	3023	44.5%	13.4%	15.8%	68.7%	13.4%	3.8%
Positive prediction fraction 25%	TP	FP	FN	TN	sensitivity	precision	MCC	AUC	AUPRC	P (sensitivity)
Full machine-learning model	140	838	89	2846	61.1%	14.3%	20.8%	75.3%	17.1%	-
Full logistic regression model	136	842	93	2842	59.4%	13.9%	19.8%	74.1%	16.7%	35.3
Parsimonious machine-learning model	134	844	95	2840	58.5%	13.7%	19.3%	74.4%	16.8%	28.3
Parsimonious logistic regression model	125	853	104	2831	54.6%	12.8%	17.0%	73.1%	16.8%	7.8
machine-learning model excluding age	121	857	108	2827	52.8%	12.4%	16.0%	72.8%	16.9%	3.6
Age-model	113	805	116	2879	49.3%	12.3%	15.2%	68.7%	13.4%	0.5
Positive prediction fraction 30%	TP	FP	FN	TN	sensitivity	precision	MCC	AUC	AUPRC	P (sensitivity)
Full machine-learning model	153	1020	76	2664	66.8%	13.0%	20.0%	75.3%	17.1%	-
Full logistic regression model	147	1026	82	2658	64.2%	12.5%	18.6%	74.1%	16.7%	27.9

Parsimonious machine-learning model	147	1026	82	2658	64.2%	12.5%	18.6%	74.4%	16.8%	27.7
Parsimonious logistic regression model	145	1028	84	2656	63.3%	12.4%	18.1%	73.1%	16.8%	21.6
machine-learning model excluding age	140	1033	89	2651	61.1%	11.9%	17.0%	72.8%	16.9%	10.2
Age-model	122	933	107	2751	53.3%	11.6%	14.8%	69.8%	13.4%	0.1

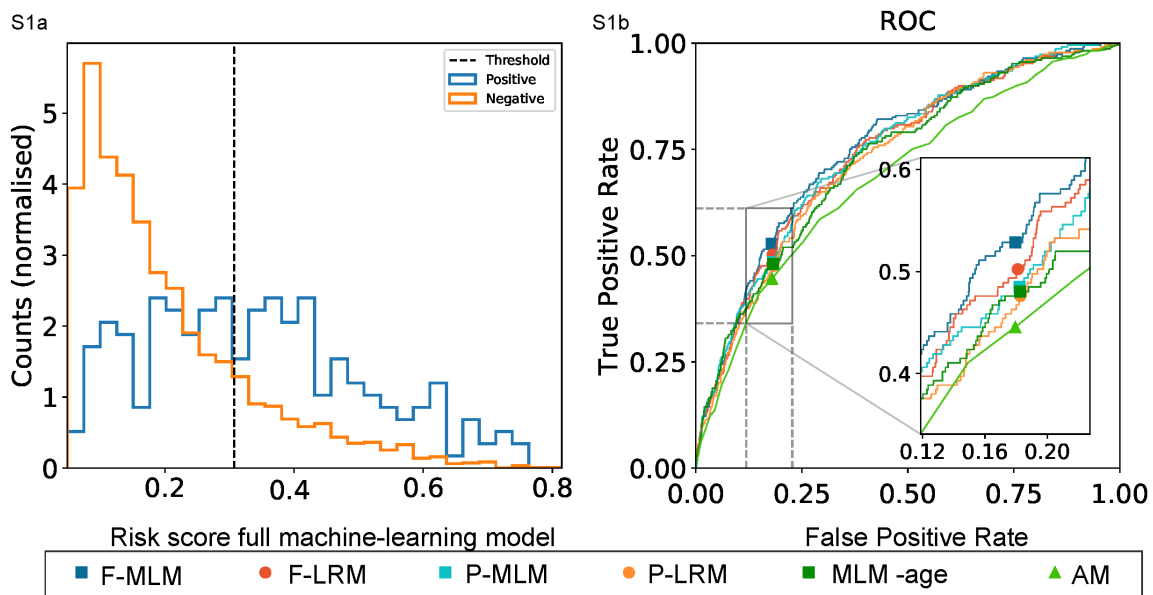
TP: true positives FP: false positives FN: false negatives TN: true negatives MCC: Matthews correlation coefficient AURC: area under the ROC curve AUPRC: area under the precision recall curve P(sensitivity): probability that the model performs better than the machine-learning model relative to sensitivity.

SDC2

[Click here to access/download;Supplemental Digital Content;SDC2.pdf](#)

## Supplemental Digital Content 2

Figure S1a-b



S1a) Distribution of full machine learning model risk scores for patients +/- outcome B(LOS >4 days or readmissions due to "medical" morbidity or LOS >4 days with no recorded morbidity). The dashed line marks the classification threshold of 20% positive prediction fraction.

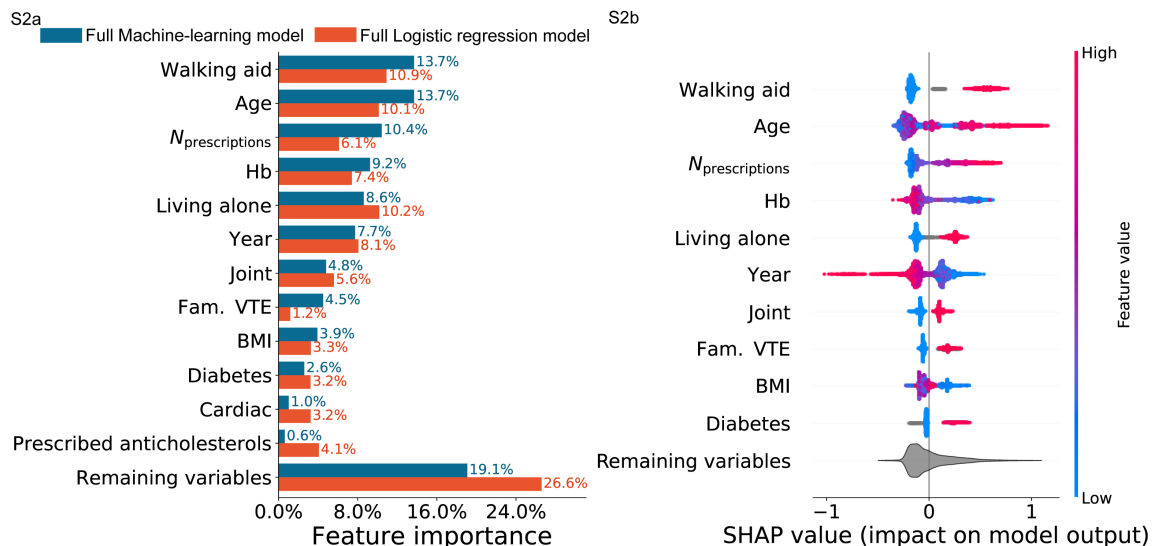
S1b) Receiver operating curves (ROC) for the full machine learning model (F-MLM), full logistic regression model (F-LRM), parsimonious machine learning model (P-MLM), parsimonious logistic regression model (P-LRM), machine learning excluding age (MLM -age) and the age-only model (AM).

SDC3

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## Supplemental Digital Content 3

Figure S2a-b



S2a) The overall importance of the 10 most important variables measured by the SHAP-values for the full machine-learning and full logistic regression models for outcome B (LOS >4 days or readmissions due to "medical" morbidity or LOS >4 days with no recorded morbidity).

Only the importance of prescribed anti-cholesterols and familiar disposition for venous thromboembolism differed between the models. The contributions of the remaining variables are summed in the bottom bar.

S2b) The SHAP-values for the full machine-learning model where values increase while negative values decrease the risk score. The color is related to the value of the variable with blue being lowest and red highest and each dot represents a patient.

## Supplemental Digital Content 4

### Figure S3a-d

SHAP scatter-plot on the contributions to the full machine-learning model on outcome B (LOS >4 days or readmission due to “medical” morbidity), for individual types of prescribed anticoagulants, cardiac drugs, psychotropics and respiratory drugs stratified by age.

#### Legend:

##### 3a) Prescribed anticoagulants

VKA: vitamin K antagonists ASA: acetylsalicylic acid DOAC: direct oral anticoagulant ADP: Adenosine diphosphate ACE: angiotensin converting enzyme

##### 3b) Prescribed cardiac drugs

ACE: angiotensin converting enzyme AHT: antihypertensive. Other AHT were defined as AHT different from diuretics ANG-II/ACE inhibitors or Ca2+antagonists. IHD: Ischemic heart disease

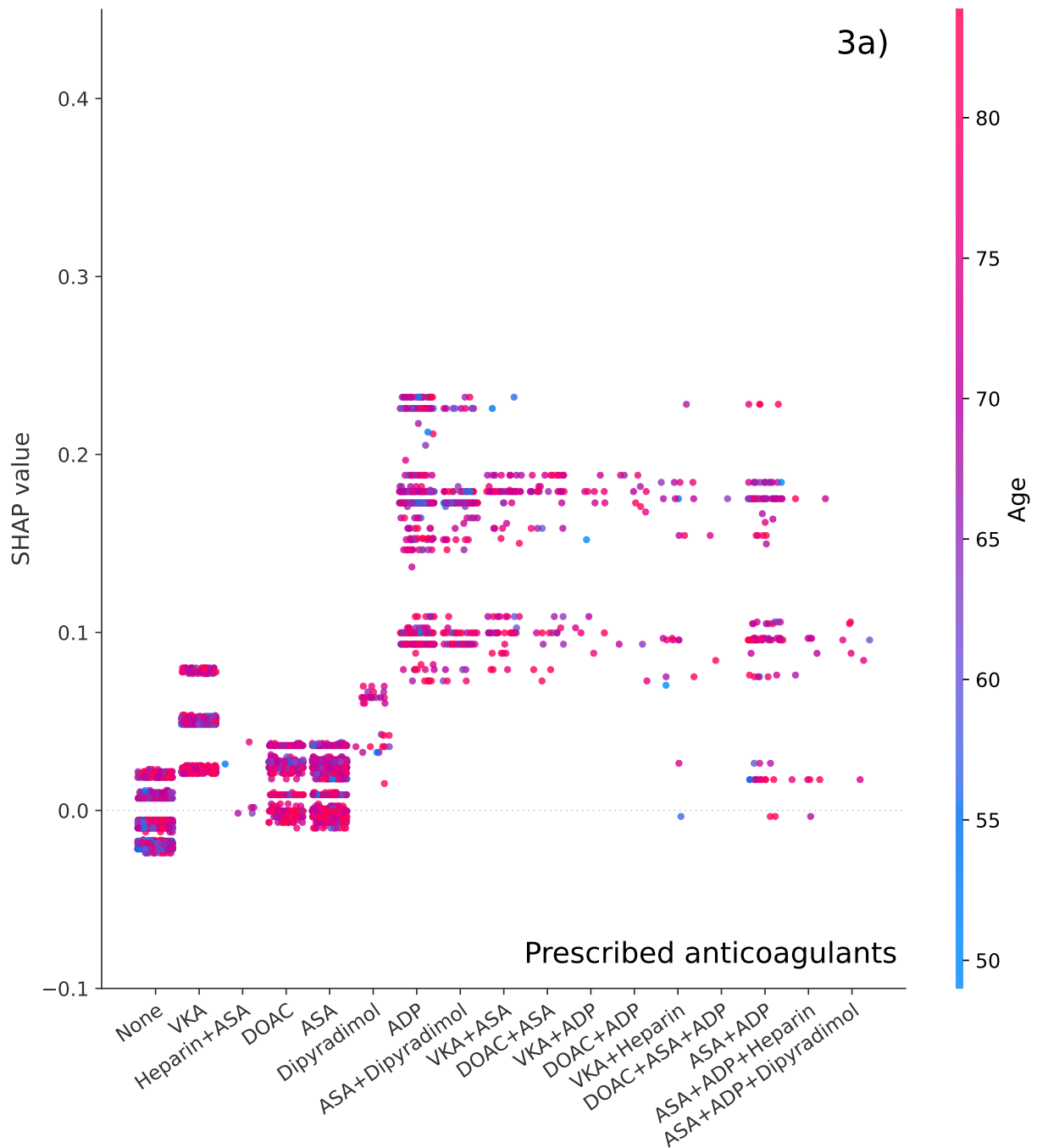
##### 3c) Prescribed psychotropics

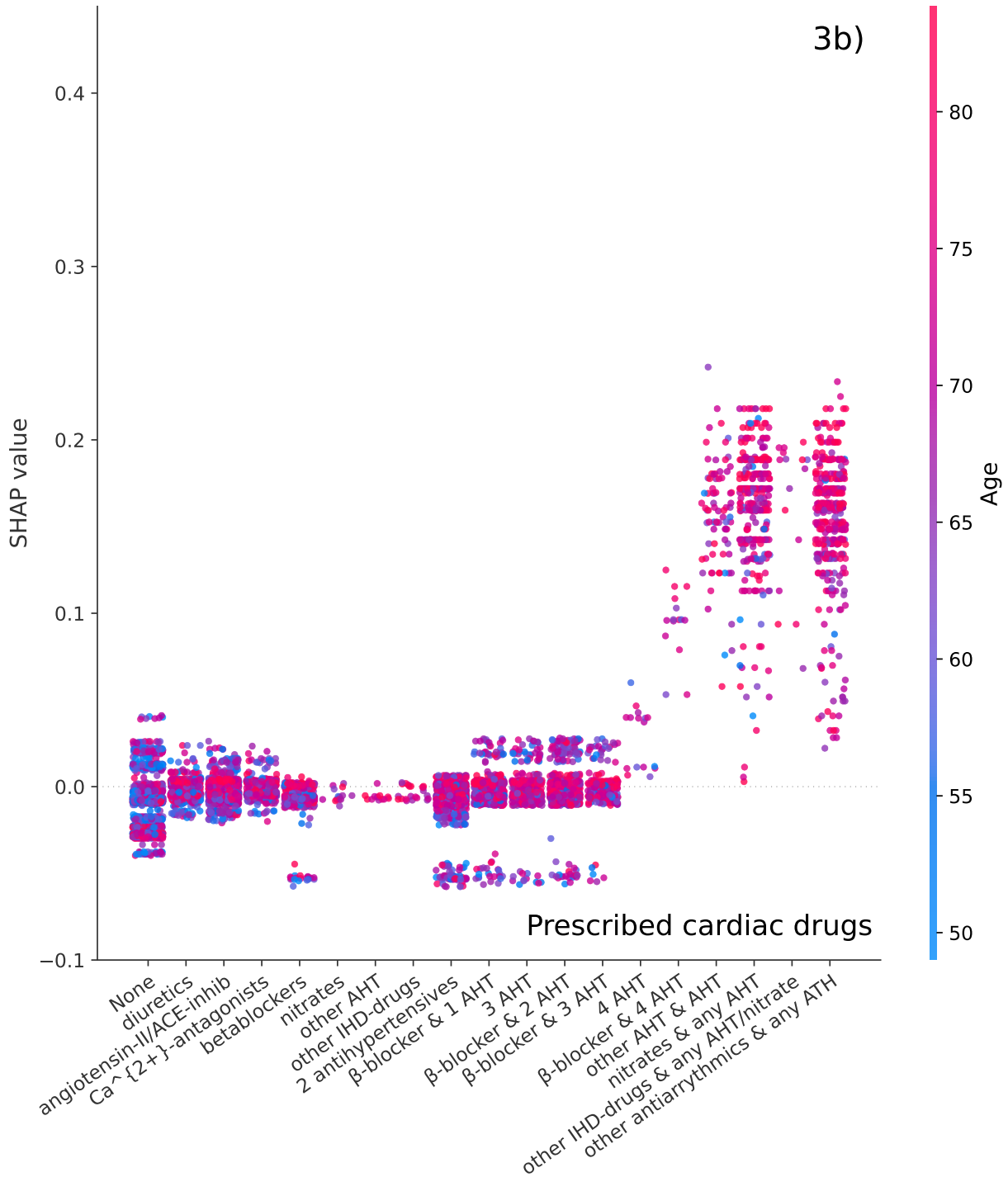
SSRI: Selective serotonin inhibitor SNRI: Serotonin and norepinephrine reuptake inhibitor NaRI: Norepinephrine reuptake inhibitor NaSSA: Norepinephrine and specific serotonergic antidepressants. AD: antidepressants BZ: Benzodiazepines (likely underreported due to limited general reimbursement in Denmark). ADHD: Attention-deficit/hyperactivity disorder

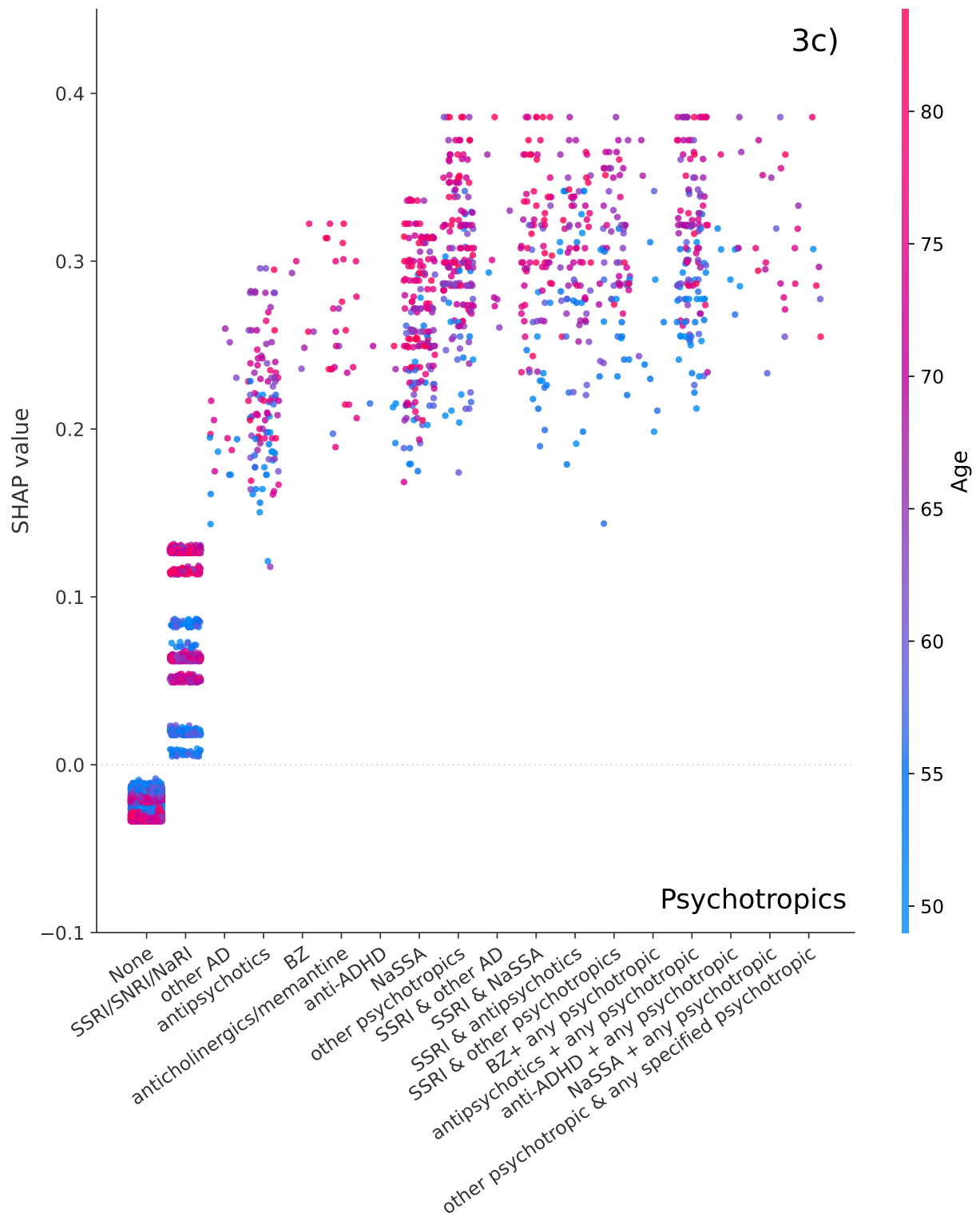
##### 3d) Prescribed respiratory drugs

The model found no additional information from this variable why all values equal 0.

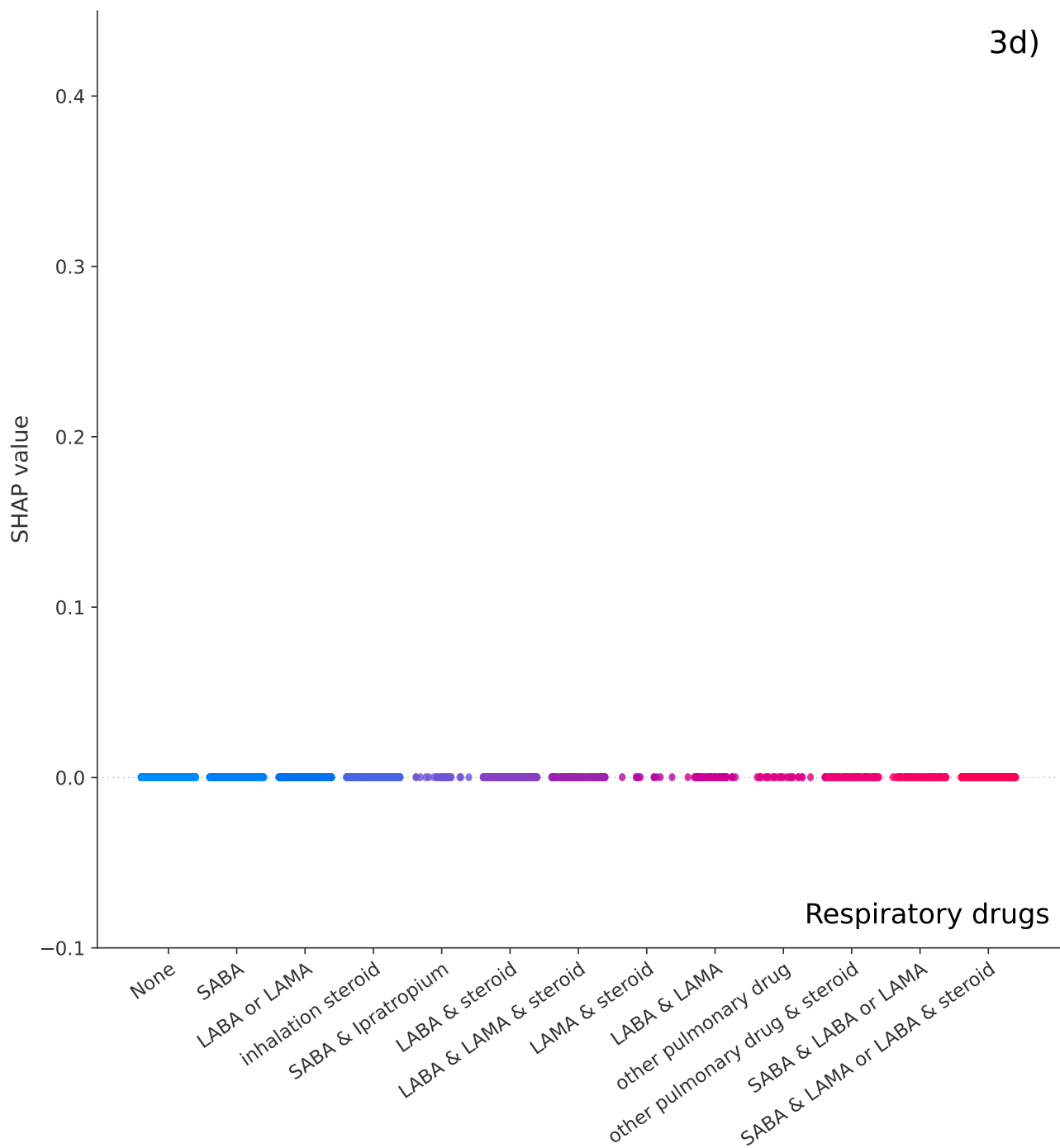
SABA: Short-acting beta agonist LABA: long-acting beta agonist LAMA: Long-acting muscarinic antagonist.







3d)





## **4 Paper III**

The following pages contain the paper:

Mathias S. Heltberg, **Christian Michelsen**, Emil S. Martiny, Lasse E. Christensen, Mogens H. Jensen, Tariq Halasa and Troels C. Petersen (2022). “Spatial Heterogeneity Affects Predictions from Early-Curve Fitting of Pandemic Outbreaks: A Case Study Using Population Data from Denmark”. In: Royal Society Open Science 9.9. issn: 2054-5703. doi: 10.1098/rsos.220018.

**Research**


**Cite this article:** Heltberg ML, Michelsen C, Martiny ES, Christensen LE, Jensen MH, Halasa T, Petersen TC. 2022 Spatial heterogeneity affects predictions from early-curve fitting of pandemic outbreaks: a case study using population data from Denmark. *R. Soc. Open Sci.* **9:** 220018. <https://doi.org/10.1098/rsos.220018>

Received: 18 January 2022

Accepted: 16 August 2022

**Subject Category:**

Mathematics

**Subject Areas:**

mathematical modelling/biophysics/  
computational biology

**Keywords:**

pandemics, agent-based modelling,  
spatial heterogeneity, fitting, COVID-19

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# Spatial heterogeneity affects predictions from early-curve fitting of pandemic outbreaks: a case study using population data from Denmark

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The modelling of pandemics has become a critical aspect in modern society. Even though artificial intelligence can help the forecast, the implementation of ordinary differential equations which estimate the time development in the number of susceptible, (exposed), infected and recovered (SIR/SEIR) individuals is still important in order to understand the stage of the pandemic. These models are based on simplified assumptions which constitute approximations, but to what extent this are erroneous is not understood since many factors can affect the development. In this paper, we introduce an agent-based model including spatial clustering and heterogeneities in connectivity and infection strength. Based on Danish population data, we estimate how this impacts the early prediction of a pandemic and compare this to the long-term development. Our results show that early phase SEIR model predictions overestimate the peak number of infected and the equilibrium level by at least a factor of two. These results are robust to variations of parameters influencing connection distances and independent of the distribution of infection rates.

<sup>†</sup>These authors contributed equally.

## 1. Introduction

Over the past years, the pathogen now known as SARS-CoV-2 has spread dramatically, risen in several waves, paralyzing societies, resulting in a large number of deaths and severe economic damage worldwide [1,2]. Mathematical models have estimated the reproduction number and guided the authorities in an attempt to minimize the damage caused by the virus [3–6]. Even though modern algorithms using machine learning have helped the process [7,8], the majority of models used to predict the size of the pandemic (or a rising wave of the disease) have been variants of the SIR/SEIR model. The SIR model was originally proposed in 1927, in the seminal work of Kermack and McKendrick, who successfully described the evolution of a pandemic, using a mean field approximation where all individuals are described as one population [9]. In the investigations of the SARS-CoV-2 pandemic, the mathematical models have varied in complexity including simple deterministic compartmental models [6,10], meta-population compartmental models [11–13], individual based models without including spatial specifications [4,14,15] and spatio-temporal agent-based models [16].

One aspect in the modelling is the ability to predict the infection peak height and the number of individuals who will be infected based on the early rise in the number of infected (before governmental interference). Earlier work has pointed out the importance of including heterogeneity when modelling the spread of infectious disease such as contact patterns between individuals [17], population mixing assumptions [18], heterogeneities caused by super-spreaders [15], and the spatial dependency of COVID-19 [19,20]. These mathematical models have not combined heterogeneous elements nor quantified how much the early SIR/SEIR predictions might be biased.

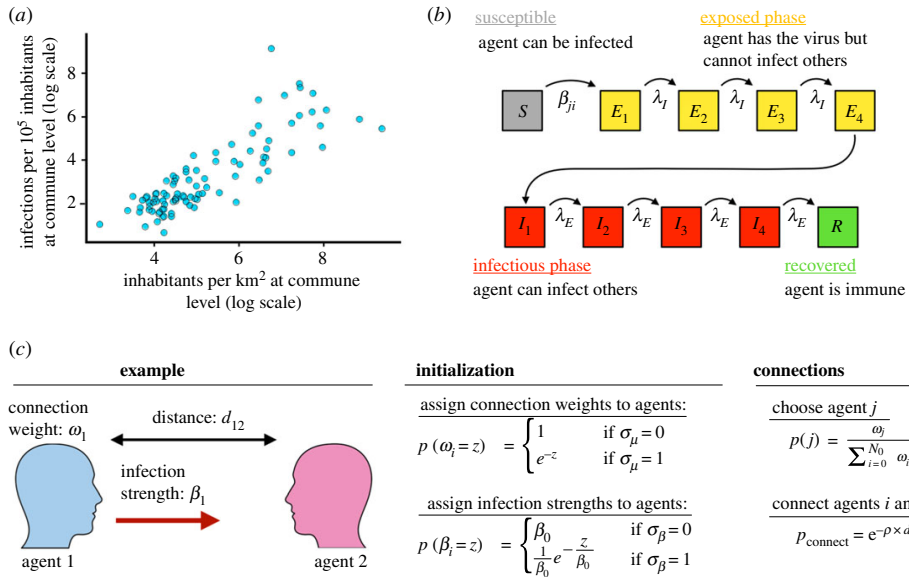
In this paper, we include geographical distributions based on an entire population, using population data of Denmark. When the SIR model was originally formulated, 95 years ago, data was not available to investigate the effects of geographical and demographic differences among the population, which might be one of the reasons why fundamental properties for diseases, such as the basic reproduction number ( $R_0$ ), can vary significantly between different regions [21]. However, with modern collection of data, these geographical aspects might be accounted for. Our main goal of this work is therefore to investigate the importance of heterogeneities in a geographically distributed population on the spread of a pandemic. We find that the heterogeneity arising from spatial inhomogeneities causes an increase in the early stage of the pandemic, affecting the initial forecast and highlighting the importance of early intervention in order to minimize the effects of the pandemic.

### 1.1. Construction of the model

In order to investigate the effect of a geographically distributed population, we extracted the number of infected per commune (from the Danish Serum Institute [22]) and divided this number with the number of inhabitants in each commune to obtain the number of infected per individual in each commune. This number we then plotted against the number of inhabitants in that specific commune (extracted from statistics Denmark [23]). Doing so, we found a strong correlation between the population density and the number of infections per inhabitant as seen in figure 1a. This observation has been made for many other countries [24–29] and underlines the aspect of disease spreading that has been observed since ancient times; that densely populated regions often have larger pandemics than the rural areas. Note that in the very early stage of a pandemic, before the exponential growth rate is reached, micro outbreaks will guide its evolution and these events can likely take place in regions with low density [30].

### 1.2. Disease simulation

To simulate evolution of the disease, we assigned each individual (agent) to a state (predominantly initialized in state S) and assigned four states to the exposed phase and four states to the infectious phase, in order to achieve an Erlang distribution (which is related to the Gamma distribution) of time in each phase [31]. Once in the exposed phase, the infected agent has a rate to move into another state, where the rate is fixed based on experimental data in order to achieve a mean time in the exposed phase of approximately 4 days (table 1). Each agent in the Infectious phase can infect other agents that have a connection to this agent in the network. This definition of agents in discrete states is naturally a simplification of the real pandemic, and we stress that this mathematical model aims at describing the spread of the disease in a simple way that does not capture all aspects of the real disease. We do not believe that this impacts our main conclusions in any way, as we are aware that one should always be careful when making these kinds of simplifications. To investigate the effect of



**Figure 1.** (a) Population density (x-axis) and the number of infections per  $10^5$  inhabitants (y-axis) for each commune in Denmark. (b) Illustration of the modified susceptible-exposed-infected-removed (SEIR) model used. It consists of 10 consecutive states ( $S, E_{1-4}, I_{1-4}$  and  $R$ ), with transition rates governed by  $\beta, \lambda_E$  and  $\lambda_I$ , respectively. (c) Illustration of how the spatial network is generated and heterogeneities in individuals included.

infection heterogeneities, we assigned an infection strength to each connection in the network, so some agents were more infectious than others. In order to control the degree of this heterogeneity, we assigned a boolean parameter  $\sigma_\beta$ , that if switched on generated an exponential distribution in infection strengths, keeping the mean field reproduction number fixed. The reproduction number between the ABM and the SIR model is related through the parameter  $\tilde{\beta} = \beta(\mu/2N_0)$ . All transitions between states and infection of other individuals were done using the Gillespie algorithm [32]. This is schematized in figure 1b.

### 1.3. Network creation

In order to construct the underlying network, we created a set-up whereby two agents were chosen at random but based on their individual connectivity weight each iteration and connected with some probability based on their spatial position. To include the possibility of highly connected individuals independent of their spatial position, we assigned a boolean parameter  $\sigma_\mu$  that, if switched on, generated an exponential distribution in weights for the individuals, keeping the mean field reproduction number fixed similar to the heterogeneity in infection strengths. To include the spatial position in the network, we introduced a parameter  $\rho$ , so the probability of connecting two chosen agents decayed exponentially with the distance between them:  $p_{\text{connect}} = e^{-\rho \times d_{ij}}$ . In order to allow some long-distance connections we introduced another parameter  $\varepsilon \in [0; 1]$ , that determines the fraction of distance-independent contacts. To construct the network of spatially distributed contacts, we chose the parameters using data based on:

- The geographical location of people in Denmark (from Boligsiden [33])
- The average number of contacts per individual per day of 11 (from HOPE [34]). Given an average infectious period of 4 days, we approximate the average number of effective contacts to be  $\mu = 40$
- The average commuting distance  $\rho = 0.1 \text{ km}^{-1}$  and the fraction of long-distance commutes  $\epsilon_\rho = 4\%$  (from statistics Denmark [23])

This is schematized in figure 1c and further described in the Methods section. All 10 parameters in this model are defined and outlined in table 1. We note that in order to keep the parameters space low, this model does not include the effects of temporal changes such as seasonality and holidays. While all agents

**Table 1.** Overview of the 10 parameters applied in this study, their typical value, and the ranges we have considered. The first six parameters are standard SEIR parameters, whereas the last four parameters define the heterogeneity in the model. These four parameters do not affect the SEIR model.

variable	description	value	range	units
$N_0$ :	population size	$5.8 \times 10^6$	$10^5 - 10^7$	—
$N_{\text{init}}$ :	number of individuals initially infected	100	$1 - 10^4$	—
$\mu$ :	average number of network contacts	40	10–100	—
$\beta$ :	typical infection strength	0.01	0.001–0.1	$d^{-1}$
$\lambda_F$ :	rate to move through $\frac{1}{4}$ of latency period	1	0.5–4	$d^{-1}$
$\lambda_I$ :	rate to move through $\frac{1}{4}$ of infectious period	1	0.5–4	$d^{-1}$
$\sigma_\mu$ :	population clustering spread	0	0–1	—
$\sigma_\beta$ :	interaction strength spread	0	0–1	—
$\rho$ :	typical acceptance distance	0.1	0–0.5	$\text{km}^{-1}$
$\epsilon_\rho$ :	fraction of distance-independent contacts	0.04	0–1	—

have been assigned parameters to their infection network that are derived from statistics of Denmark for both employees and students, we have not divided each agent into specific occupations.

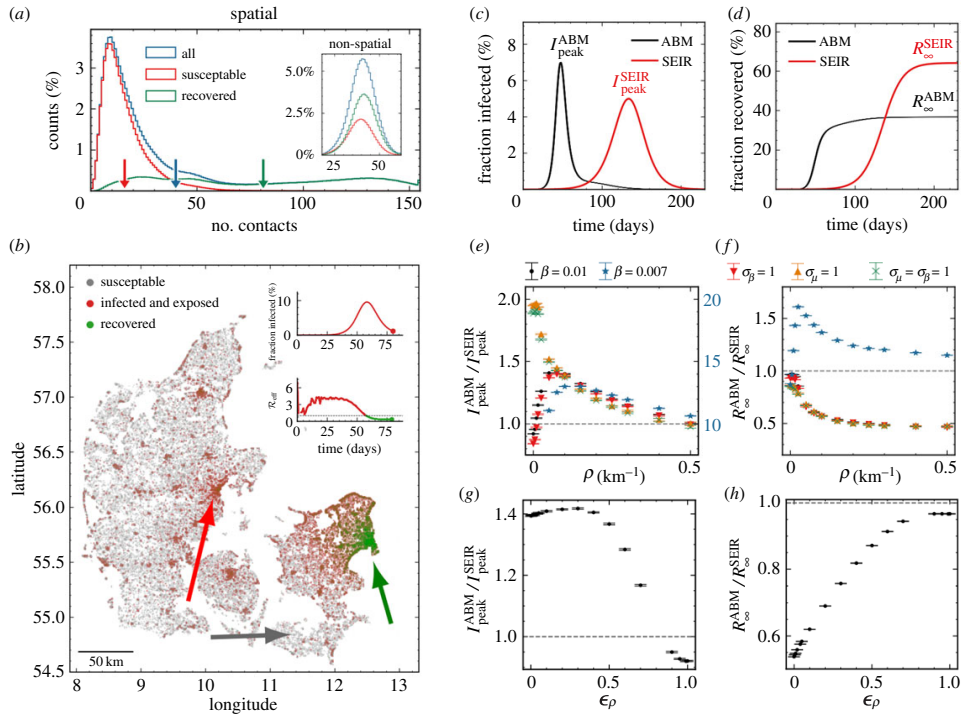
Before including heterogeneity, we compared the ABM to the corresponding SEIR model as a test, and found them to agree within 5% for all parameter configurations tested. Here, we also tested the effect of the number of individuals initially infected (see electronic supplementary material). This concludes that the SEIR and ABM model are calibrated to have the same reproduction number in the absence of heterogeneities. Next, we will introduce heterogeneities into the system, while keeping the sum of contacts and infection strengths constant, to study how this affects the evolution of the pandemic.

## 2. Results

### 2.1. Geographical distributions in a population and large variances in numbers of contacts leads to increased infection levels

Having introduced heterogeneity, the distributions of connections in this network were created automatically through the population clustering, see figure 2a. This naturally leads to individuals living in densely populated areas having higher number of connections. In an example simulation with 100 initially infected individuals,  $N_{\text{init}} = 100$ , we observed a spatial difference in areas affected by the disease (figure 2b), as expected. Note that we also show the effective reproduction number ( $\mathcal{R}_{\text{eff}}$ ) as a function of time in the lower part of the inserted panel. One region reached local endemic steady state (green arrow, figure 2b) while other regions of similar density were highly infected (red arrow, figure 2b) and yet other districts were almost unaffected (grey arrow, figure 2b). To quantify the effect of population clustering, we compared the ABM result to the reference SEIR model of similar parameters. Generally, we observed that the epidemic developed faster with a higher infection peak  $I_{\text{peak}}$ , but also subsided quicker, leading to a lower number of infected once reaching endemic steady state,  $R_\infty$  (figure 2c,d).

In order to explore how population clustering affects the epidemic, we chose a reference value of infection rates,  $\beta = 0.01$ , and an alternative value of  $\beta = 0.007$ . In the absence of spatial dependence ( $\rho = 0 \text{ km}^{-1}$ ), these correspond to initial reproduction numbers  $\mathcal{R}_0 \approx 1.7$  and 1.1, respectively. Here, we define the reproduction number as the average number of agents each infectious agent will infect in the first part of the disease. Increasing the spatial dependence (i.e. increasing  $\rho$ ) leads to a significant rise in the infection peak for the ABM,  $I_{\text{peak}}^{\text{ABM}}$ , compared to the (unaffected) SEIR model,  $I_{\text{peak}}^{\text{SEIR}}$  for both the reference value and the alternative lower value of  $\beta$  (black and blue points, figure 2e). We introduced heterogeneity in infection strengths ( $\sigma_\beta = 1$ , see figure 1b), thus making some individuals much more infectious than others (i.e. including *super shedders*). We found no significant impact from this effect (red points in figure 2e). Similarly, we introduced heterogeneity in connection weights ( $\sigma_\mu = 1$ , see figure 1b), thus making some individuals much more likely to form contacts than others



**Figure 2.** (a) Histograms showing the number of susceptible (red) and recovered (green) individuals at the end of an epidemic with  $\rho = 0.1 \text{ km}^{-1}$ . The distribution before the epidemic is shown in blue. The arrows show the mean of each distribution. The inset shows the same for  $\rho = 0 \text{ km}^{-1}$ . (b) Visualization of the spatial position of individuals during the infection and which state they are in. Green arrow: largest city in Denmark (Copenhagen): mostly recovered. Red arrow: Second largest city in Denmark (Aarhus): mostly infected. Grey arrow: low-population area: mostly susceptible (i.e. have not been infected). (c) Number of infected individuals as a function of time. Data shown for the spatially distributed network ( $\rho = 0.1 \text{ km}^{-1}$ ). Simulation was repeated 10 times. (d) Cumulative sum of individuals who have had the disease as a function of time (with  $\rho = 0.1 \text{ km}^{-1}$ ). (e) Relative difference in maximal number of infected,  $I_{\text{peak}}$ , between deterministic (SEIR) and ABM as a function of  $\rho$ , and shown for different parameters. Note the data for  $\beta = 0.007$  are shown in blue with a factor 10 scaling (right y-axis). (f) Relative difference in total number of infected at the end of the epidemic,  $R_\infty$ , between deterministic (SEIR) and ABM as a function of  $\rho$ . Colours similar to (e). (g) Same as (e), but as a function of  $\epsilon_\rho$ . (h) Same as (f), but as a function of  $\epsilon_\rho$ .

(i.e. including *super connecters*). This leads to a significant effect for  $\rho = 0 \text{ km}^{-1}$ , which converges towards the other curves for  $\rho > 0.1 \text{ km}^{-1}$  (orange (only super connecters) and green (super connecters and super shedders) points in figure 2e). The total number of individuals that have been in the infectious state, when there are not enough susceptible agents for the disease to keep infecting new individuals, is termed  $R_\infty$ , and this converged towards half of the SEIR model prediction as a function of  $\rho$  except for  $\beta = 0.007$  where the endemic steady state level is larger than the one obtained by the SEIR model (figure 2f). We note that in reality, individuals can lose immunity and therefore new waves can emerge. But for a completely susceptible population,  $R_\infty$  describes the fraction of the population that will get the disease during a specific wave. Fixing  $\rho = 0.1 \text{ km}^{-1}$  and increasing the fraction of distance-independent contacts,  $\epsilon_\rho$ , we found that  $I_{\text{peak}}^{\text{ABM}}$  is almost unaffected for  $\epsilon_\rho < 0.5$  (figure 2g), while  $R_\infty^{\text{ABM}}$  increases linearly towards the SIER model  $R_\infty^{\text{SEIR}}$ , as expected (figure 2h).

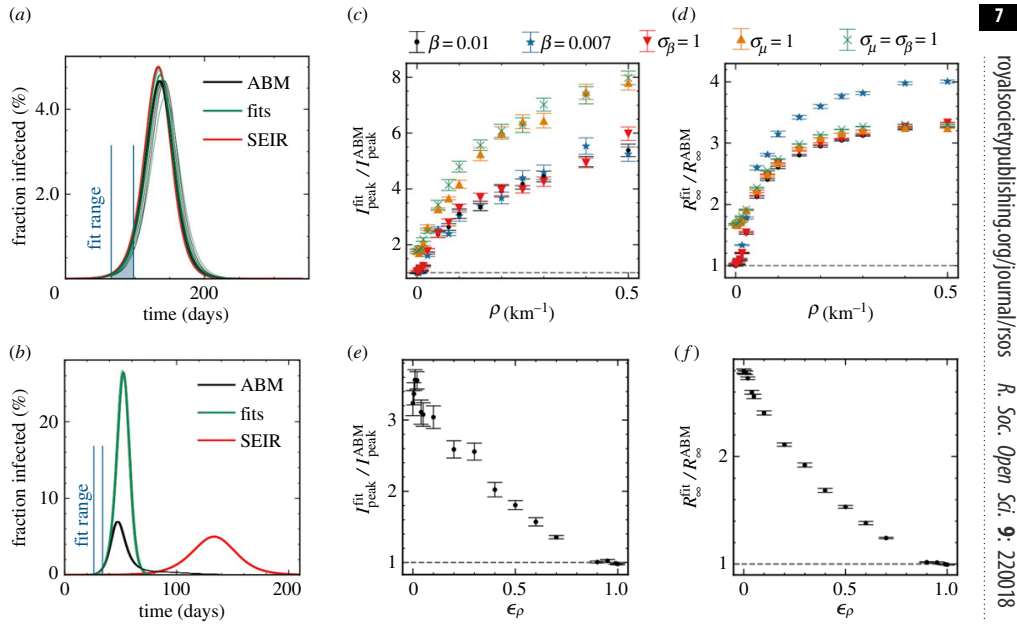
## 2.2. Fitting early infection curves leads to significant bias in estimating the size of the pandemic

Next, we consider how these heterogeneities bias the traditional SEIR model predictions, especially the predictions based on fits to the number of infected (i.e. the curve to be flattened) in the beginning of the epidemic (see Methods). Without spatial dependence, the predicted curves fitted the number of infected

individuals very well (figure 3a). Introducing spatial dependence ( $\rho = 0.1 \text{ km}^{-1}$ ) leads to a severe overestimation of the epidemic based on the number of early infection cases (figure 3b). This result can be interpreted by the fact that in societies where population density and thus individual contact number varies significantly, the early phase will be driven by people with many contacts (*super connectors*). This typically happens in cities where the population density is high. Increasing the spatial dependence  $\rho$ , we found that the SEIR model predictions overestimated the infection peak height  $I_{\text{peak}}$  and the total number of infected  $R_{\infty}$  significantly even for very small spatial heterogeneities (figure 3c, d). We observed this general trend for all tested combinations of parameters and heterogeneities. In particular, we found that if long-distance connections  $\epsilon_{\rho}$  are below 10%, the bias in the estimated infection peak height,  $I_{\text{peak}}$ , was constant within statistical uncertainty (figure 3e). For the total number of infected,  $R_{\infty}$ , we observed an almost linear regression to the SEIR model as  $\epsilon_{\rho}$  approaches one. However, even when  $\epsilon_{\rho} = 0.25$ , the prediction bias was still a factor of two (figure 3f). We concluded from these curves a general trend; if one fits an SEIR model to infection numbers during the beginning of an epidemic, and use these estimates to predict the characteristics of the epidemic at a national level, one overestimates the number of infected by at least a factor of two.

### 3. Discussion

In summary, this work outlines that the degree of population clustering in Denmark creates a discrepancy between the early predictions made by the SEIR models and the underlying agent-based interactions. It results in a significant overestimation of the impact of the disease, both in terms of maximal number of simultaneously infected (by a factor of 3) and the endemic steady state level (by a factor of 2.5). Such discrepancies have been observed for earlier pandemics, for instance, the 1918 Spanish flu, where the predicted number of individuals that would get the disease within a season turned out to be higher than the actual outcome [35]. The present results can be an important element in explaining these mismatches, even though other elements, such as for instance social distancing and the population behaviour, play a vital part. When facing a rising pandemic, societies are faced with the task of laying out strategies to minimize the consequences, including the importance of *flattening the curve*. While this is truly crucial to avoid overpopulated hospitals, the understanding of the pandemic should be taken seriously enough that we might specify to a higher degree of certainty which curve to be flattened. Our results highlight an important element in the prediction of infection levels and quantify the effect of density heterogeneities. We are aware that these results are not directly applicable to the pandemic of SARS-CoV-2 as a whole, since numerous mutations have increased the infection rates compared to the early estimates and created a strong heterogeneity in the infection worldwide. Furthermore, the actual evolution of the pandemic was highly affected by the different governmental interventions, that are not included in this work. However, this study emphasizes the abnormally large reproduction rates in the beginning of a pandemic, due to individuals with more connections than the rest of the population and attempts to quantify this bias, when countries should estimate the severity of a disease based on the data collected in the early phase. This also underlines the benefits by making lockdowns early in the pandemic, when a population is highly susceptible (for instance to a new mutation) and therefore can be driven by *super connectors*. Since people living in city-clusters are more likely to have many contacts, or infection events, they are on average more likely to be affected in the early stage of the pandemic (if they do not implement social distancing). By removing contacts from these individuals, through some level of interaction in order to reduce the number of social contacts, one can avoid the worst peak while affecting the lowest number of people. While our work describes some fundamental aspects of the disease spreading, this model does not consider asymptomatic individuals, which has been an important aspect of the SARS-CoV-2 pandemic [36,37]. Effectively, asymptotic individuals would correspond to a very heterogeneous distribution of time the agents spend in the infectious state. While agents with symptoms would predominantly isolate themselves and thereby significantly reduce their ability to infect other agents, asymptomatic agents would have a long time in the infectious state, thereby infecting more individuals. In this work, we have not considered the observation that individuals lose their immunity to SARS-CoV-2 which was first studied in the Brazilian city of Manaus. For this model, the temporal decline of immunity would lead to more pandemic ‘waves’, but for a fixed disease transmissibility this would not alter the maximal height of the peak number of infected, since this occurred for all the initially susceptible population. Finally, we note that this work does not include a vast range of divisions for the population, including age, socio-economic status etc. We have not included this directly, since we wanted to estimate as cleanly as possible how the heterogeneity in the



**Figure 3.** (a) Number of infected individuals for the ABM in black, the SEIR model in red and the SEIR fits to the ABM data in green. Blue lines show the interval where parameters are fitted (also shown below the curves). Here,  $\rho = 0 \text{ km}^{-1}$ . (b) Same as (a) but with population clustering ( $\rho = 0.1 \text{ km}^{-1}$ ). (c) Relative difference in maximal number of infected,  $I_{\text{peak}}$ , between the fit and the ABM for different values of  $\rho$ . Simulations repeated 10 times for each data-point. (d) Relative difference in total number of infected at the end of the epidemic,  $R_{\infty}$ , between the fit and the ABM for different values of  $\rho$ . (e) Same as (c), but as a function of  $\epsilon_{\rho}$ . (f) Same as (d), but as a function of  $\epsilon_{\rho}$ .

contact pattern, arising from a geographically distributed population, could affect the evolution of a disease. We are aware that for instance the distribution of age has an enormous impact on the health risk and that this risk is vital in the prediction of hospitalizations in modern society. However, our aim was to understand the bias in the prediction of a disease, based on the data that comes during the early periods of a disease, independently of the mortality of this disease. Mathematical predictions of disease progression have been heavily criticized [38,39] and it is important to improve the theoretical foundations of the mathematical descriptions, in order to increase the confidence in the predictions. Our work highlights the importance of estimating the spatial clustering and connectivity skewness in the population in order to correct the predictions based on SEIR models, by quantifying their biases from not including spatial clustering. We hope that this work could serve as an input to the modelling and prediction of future pandemics and the importance of avoiding super-spreaders in high-density areas.

### 3.1. Methods

#### 3.1.1. Construction of spatial network

We initialized  $N_0$  agents on a network generating a total of  $\mu \times N_0$  links between two agents, with an assigned interaction strength  $\beta_{ij}$  for each link. The average contact number,  $\mu$ , was fixed to 20, based on results from the Danish HOPE project, gathering data on population behaviour since April 2020 [34]. In order to include a realistic, geographical distribution of the population, we randomly selected agent locations from a two-dimensional kernel density estimate we had generated based on housing sales in Denmark 2007–2019 (data given with permission from Boligsiden, [33]). We note that in this distribution, we do not take specific geographical elements such as roads or environment into account (which has been previously studied for other diseases [40]) as we assume that this effect is small in a country like Denmark, where all parts are connected and natural obstacles such as mountains and rivers are not present. To connect the agents, we used a hit and miss method, where two random agents are first suggested and then connected with probability,  $p(d) = e^{-\rho d_{ij}}$ . Here,  $d_{ij}$  is the distance between agents and

$\rho$  is a parameter with units of inverse distance. We choose  $\rho = 0.1 \text{ km}^{-1}$  (i.e. 10 km) which is the average distance travelled by labour force (statistics Denmark [23]). To allow some long-distance interactions, we introduced a parameter  $\epsilon_\rho = 4\%$  representing the fraction of distance-independent connections. This value is based on the fraction of workers travelling longer than 50 km to work (statistics Denmark [23]).

### 3.1.2. Fits and predictions

We defined an early phase to be the period of time when between 0.1% and 1% of the population were infected (blue lines figure 3a). We then fitted  $\beta$  and a time delay,  $\tau$ , to the SEIR model with a  $\chi^2$ -fit (assuming Poissonian statistics) and kept  $\lambda_E$  and  $\lambda_I$  fixed to the true numbers (used in the simulation). The initial number of infected,  $N_{\text{init}}$ , was also fixed to the true numbers. The fit parameters were then inserted into the SEIR model, and  $I_{\text{peak}}^{\text{fit}}$  and  $R_{\infty}^{\text{fit}}$  were extracted from the fitted model and compared to the  $I_{\text{peak}}^{\text{ABM}}$  and  $R_{\infty}^{\text{ABM}}$  from the ABM simulation.

Data accessibility. Data and relevant code for this research work are stored in GitHub: [www.github.com/ChristianMichelsen/NetworkSIR](https://www.github.com/ChristianMichelsen/NetworkSIR) and have been archived within the Zenodo repository: <https://zenodo.org/badge/latestdoi/258223118>.

Authors' contributions. C.M.: conceptualization, formal analysis, investigation, methodology, software, validation, visualization, writing—original draft, writing—review and editing; E.S.M.: investigation, software, validation, visualization, writing—review and editing; L.E.C.: supervision, validation, writing—review and editing; T.C.P.: conceptualization, investigation, methodology, project administration, software, supervision, validation, visualization, writing—original draft, writing—review and editing; M.L.H.: conceptualization, formal analysis, investigation, methodology, software, validation, visualization, writing—original draft, writing—review and editing; M.H.J.: formal analysis, investigation, supervision, validation, writing—review and editing; T.H.: conceptualization, investigation, supervision, validation, visualization, writing—review and editing.

All authors gave final approval for publication and agreed to be held accountable for the work performed therein. Conflict of interest declaration. We declare that we have no competing interests.

Funding. M.L.H. acknowledges the Carlsberg Foundation grant no. CF20-0621 and the Lundbeck Foundation grant no. R347-2020-2250. E.S.M. and M.H.J. acknowledge support from the Independent Research Fund Denmark grant no. 9040-00116B and Danish National Research Foundation through StemPhys Center of Excellence, grant no. DNRF116. Acknowledgements. The authors are grateful to the Danish expert group of SARS-CoV-2 modelling led by Statens Serum Institute, especially Robert L. Skov, Kåre Mølbak, Camilla Holten Møller, Viggo Andreasen, Kaare Græsbøl, Theis Lange, Carsten Kirkeby, Frederik P. Lyngse, Matt Denwood, Jonas Juul, Sune Lehman, Uffe Thygesen and Laust Hvas Mortensen. Furthermore, we thank Kim Sneppen for valuable discussions.

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## **5 Paper IV**

The following pages contain the paper:

Susmita Sridar, Mathias S. Heltberg, **Christian Michelsen**, Judith M. Hattab, Angela Taddei (2022). “Microscopic single molecule dynamics suggest underlying physical properties of the silencing foci”.

# Microscopic single molecule dynamics suggest underlying physical properties of the silencing foci

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## Abstract

In order to obtain fine-tuned regulation of protein production while maintaining cell integrity, it is of fundamental importance to living organisms to express a specific subset of the genes available in the genome. One way to achieve this is through the formation of subcompartments in the nucleus, known as foci, that can form at various locations on the DNA fibers and repress the transcriptional activity of all genes covered. In this work we investigate the physical nature of such foci, by applying single molecule microscopy in living cells. Here we study the motion of the protein SIR3. By combining various statistical methods, and combining a frequentist with a bayesian approach, we extract the diffusion properties for motion in a repair foci. In order to obtain useful information based on this, we derive similar measures for the foci itself, the motion of SIR3 outside the foci and other mutants of the cell. We reveal that the behaviour inside a repair foci is highly immobile and we compare this to theoretical expressions. Based on this we hypothesize that the repair foci is probably not a result of a second order liquid-liquid phase separation but rather a so-called Polymer Bridgng Model with numerous binding sites.

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**Data availability:** Data availability is available on Zenodo or the Github repository.

**Funding:** This work was supported by XXX Foundation. The funders had no role in the decision to publish.

**Competing interests:** The author declare no competing interests.

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## 1 | INTRODUCTION

- 26 Understanding the physical principles of how cells can express and silence specific regions of the genome presents one of the most fundamental challenges in biology. As a model to study this,
- 28 budding yeast chromosomes is a strong candidate, since it has very few repetitive sequences outside of the rDNA compared to other eukaryotes that contain centromeric heterochromatin. When
- 30 haploid cells grow at their maximal rate, one characteristic aspect is that 32 telomeres accumulate at the nuclear envelope allowing them to form  $\approx$ 3–5 foci. The sizes of these are in the order of a few
- 32 hundreds of nanometer and therefore below the diffraction limit of conventional epifluorescence microscopes.
- 34 Inside such foci, the silent regulatory factors Sir2, Sir3 and Sir4 concentrate into the form of the SIR complex (Palladino et al., 1993). These are therefore termed silencing foci, since they can re-
- 36 press the expression of the underlying genes through interaction with the telomeric protein Rap1, and thereby spread on chromatin and potentially forming a compact chromatin structure. Studies
- 38 in vitro has revealed that this complex associates with nucleosome in a 1:2:1 stoichiometry and can significantly compact chromatin (Swygert et al., 2018).
- 40 The sequestration of SIR proteins from silent chromatin favor the subtelomeric repression and the position of telomeres inside these foci favors faithful recombination events upon double strand
- 42 break (Batté et al., 2017). Furthermore, it also prevents the binding of the SIRs at specific groups of promoters in the genome (Maillet et al., 1996; Marcand et al., 1996; Taddei et al., 2009).
- 44 In the foci, the telomere composition is not fixed, however telomeres show preferential attachment to other telomeres coupled to chromosome arms of approximately equal length (Therizols
- 46 et al., 2010; Schober et al., 2008; Duan et al., 2010). This process of telomeres grouping in a limited number of foci requires Sir3 association to telomeres but is independent of heterochromatin
- 48 formation (Ruault et al., 2011) and these foci has been revealed to fuse into bigger foci or hyper-clusters when SIR3 is overexpressed, suggesting a regulatory role on telomere clustering for SIR3
- 50 (Ruault et al., 2011).
- In this work we investigate the physical mechanism of the formation of silencing foci. In particular
- 52 we use using Single Particle Tracking (SPT) and Photo Activable Localization Microscopy (PALM) in

Saccharomyces cerevisiae cells in order to obtain precise information about the dynamics of single particles in the heterogeneous environment. In this, SPT is a powerful technique that makes the microscopic steps taken by the molecules observable, by taking “live” recordings of individual molecules in a cell at high temporal and spatial resolution (50 Hz, 30 nm) (Dolgin, 2019; Manley et al., 2008; Oswald et al., 2014). Based on this in vivo movement, SPT allows for grouping specific proteins into subpopulations defined by the measured diffusion coefficients. From this it is possible to quantify the motion of each subpopulation and thereby estimating the residence times in different parts of the nucleus, allowing us to estimate the free-energy of the system. To assist the SPT measurements, PALM can establish a density maps of the molecules of interest by their position at 30 nm resolution.

Using these methods we have assessed the dynamics of SIR3 cells with silencing foci. We find that inside the silencing foci, SIR3 moves significantly slower and we relate this to the motion of the whole focus itself. This allows us to identify the diffusion properties of both free telomeres, and telomeres inside a focus. Next we apply Sir4 deprived mutants and observe that the foci has disappeared, allowing us to extract the free diffusion coefficient of SIR3. Finally we use this to extract the free energy of the molecules inside the repair foci, and we compare this to the theoretical prediction, assuming that the repair foci belongs to the Polymer-Bridging model. Here we find a good agreement, thus suggesting that the physical nature of these foci is really a dense collection of multiple binding sites that suppress the movement of molecules while enhancing their concentration is the formed region.

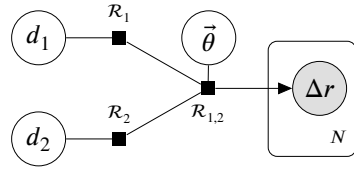
## 2 | METHODS & MATERIALS

### 2.1 | Diffusion model

For each of the different types of data (XXX), we load in the cells and group them by cell number and ID. For each group we compute the distance  $\Delta r$  between the subsequent observations  $\vec{x}_i$ :

$$\Delta r_i = \|\vec{x}_{i+1} - \vec{x}_i\|. \quad (1)$$

E.g., for Wild Type 1, we find 914 groups across 43 different cells, leading to a total of  $N = 10.025$  distances. We model the diffusion distances with a Rayleigh likelihood, where the Rayleigh distri-



**Figure 1.** A graphical representation of the Bayesian model case of two diffusion components using the directed factor graph notation (Dietz, 2022). Here  $d_1$  is the diffusion coefficient,  $\mathcal{R}_1$  is the  $d$ -parameterized Rayleigh distribution and  $\mathcal{R}_{1,2}$  is the mixture model of the Rayleigh distributions with a  $\theta$  prior.

80 bution is given by:

$$\text{Rayleigh}(r; \sigma) = \frac{r}{\sigma^2} e^{-r^2/(2\sigma^2)}, \quad r > 0. \quad (2)$$

82 In this study, we parameterize the Rayleigh distribution in terms of the diffusion coefficient  $d$ , which is related to the scale parameter  $\sigma$  in eq. (2), through the XXX parameter,  $\tau$ :

$$84 \quad \sigma = \sqrt{2d\tau}, \quad (3)$$

with  $\tau = 0.02$  in the current study. In the simplest form, where we assume only a single diffusion

86 coefficient,  $d$ , the Bayesian model for this process is:

[d prior]	$d \sim \text{Exponential}(0.1)$
88 [transformation]	$\sigma = \sqrt{2d\tau}$
90 [likelihood]	$\Delta r_i \sim \text{Rayleigh}(\sigma).$

(4)

A more realistic diffusion model include more than a single diffusion coefficient. Figure 1 shows 92 this for the two-component case in directed factor graph notation (Dietz, 2022). In particular, the figure shows the combination of the  $K = 2$  diffusion coefficients  $d_k$  through a mixture model  $\mathcal{R}_{1,2}$  of 94 the two  $d$ -parameterized Rayleigh distributions  $\mathcal{R}_k$  with a  $v$ -prior. We model each of the distances as independent, indicated by the  $N$ -replications plate. In equations, the figure is similar to:

96 [d1 prior]	$d_1 \sim \text{Exponential}(0.1)$
[d2 prior (ordered)]	$d_2 \sim \text{Exponential}(0.1), \quad d_1 < d_2$
98 [theta-hat prior]	$\theta \sim \text{Uniform}(0, 1), \quad \vec{\theta} = [\theta_1, 1 - \theta_1]$
[mixture model]	$\mathcal{R}_{1,2}(d_1, d_2, \vec{\theta}) = \text{MixtureModel}\left([\mathcal{R}(d_1), \mathcal{R}(d_2)], \vec{\theta}\right)$
100 [likelihood]	$\Delta r_i \sim \mathcal{R}_{1,2}(d_1, d_2, \vec{\theta}).$

(5)

## 102 2.2 | Model comparison

<sup>1</sup> ordered such that  $d_1 < d_k < d_K$  to prevent the classical label-switching problem in the case of mixture models (McLachlan and Peel, 2004)

We can generalize the  $K = 2$  diffusion model to higher values of  $K$  by having  $d_1, \dots, d_K$  ordered<sup>1</sup> diffusion coefficients and letting the mixture model's  $\bar{\theta}$ -prior be a random variable from a flat Dirichlet distribution (such that  $\sum_k \theta_k = 1$ ). We find that including up to three diffusion coefficients yields appropriate results. To compare the three models of different complexity, we compute the Widely Applicable Information Criterion (WAIC) (Watanabe, 2010) which is a generalized version of the Akaike information criterion (AIC) useful for Bayesian model comparison (Gelman, Hwang, and Vehtari, 2014). In short, the WAIC is an approximation of the out-of-sample performance of the model and consists of two terms, the log-pointwise-predictive-density, lppd, and the effective number of parameters  $p_{\text{WAIC}}$ :

$$112 \quad \text{WAIC} = -2 (\text{lppd} - p_{\text{WAIC}}). \quad (6)$$

The lppd is the Bayesian version of the accuracy of the model and  $p_{\text{WAIC}}$  is a penalty term related to the risk of over-fitting; complex models (usually) have higher values of  $p_{\text{WAIC}}$  than simple models, (McElreath, 2020). The minus 2 factor is just a scaling included for historical reasons leading to low WAICs being better. Given two models, A and B, we compute both the individual WAIC values,  $W_A$  and  $W_B$ , their standard deviations,  $\sigma_{W_A}$  and  $\sigma_{W_B}$ , their difference,  $\Delta_{A,B}$ , and the standard error of their difference,  $\sigma_{\Delta_{A,B}}$ .

## 112 2.3 | Implementation

120 The data analysis has been carried out in Julia (Bezanson et al., 2017) and the Bayesian models  
 121 are computed using the Turing.jl package (Ge, Xu, and Ghahramani, 2018). We use Hamiltonian  
 122 Monte Carlo sampling (Betancourt, 2018) with the NUTS algorithm (Hoffman and Gelman, 2011).  
 123 In particular, each Bayesian model have been run with 4 chains, each chain 1000 iterations long  
 124 after discarding the initial 1000 samples ("warm up").

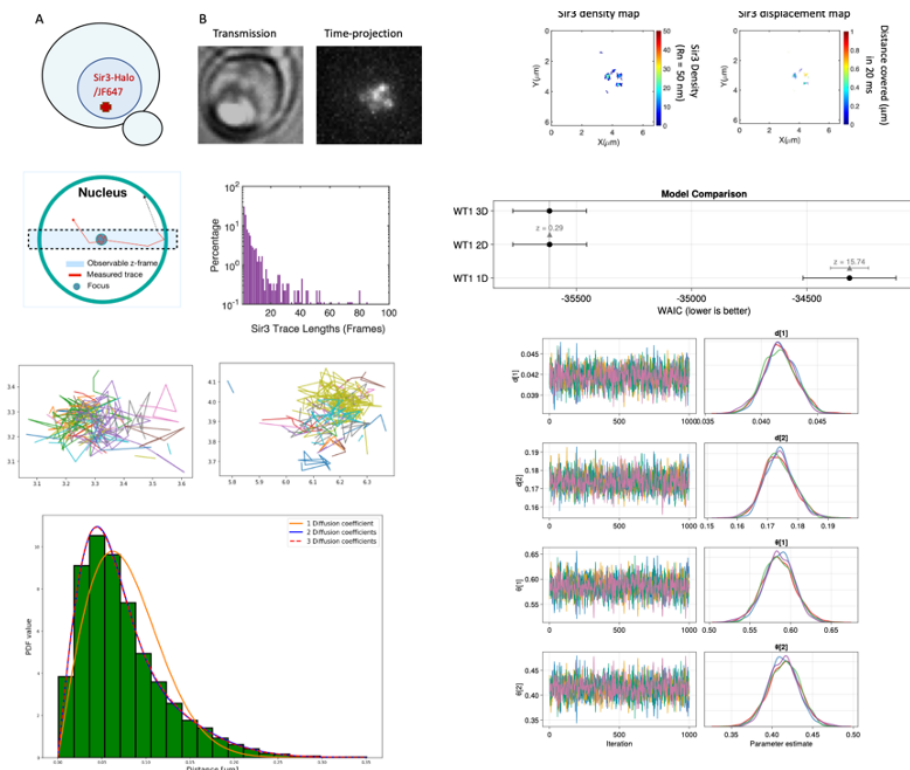
## 3 | RESULTS

### 126 3.1 | Two diffusive populations identified at for SIR3 mobility in WT

We started out by using SPT to investigate the mobility of individual SIR3 proteins in vivo in WT  
 127 cells. To obtain this imaging of SIR3 without altering its normal expression level, we constructed  
 128 a line of haploid cells that express the endogenous SIR3 fused to Halo (Figure 2A and Materials

and methods). Before we visualized this on a PALM microscope (see Materials and methods), we incubated the exponentially growing cells with fluorescent and fluorogenic JF647. This is a dye that emits light when it is bound to Halo. We then used a low concentration of JF647 in order to obtain visible individual molecules (Ranjan et al., 2020; Figure 1B). With this setup, the SIR3-Halo bound to JF647 (SIR3-Halo/JF647) were visualized at 20 ms time intervals (50 Hz) in 2-dimensions during 1000 frames until all signal had decayed. A typical individual cell is shown in Figure 2B and the tracking of the individual molecules is visualised in Figure 2C and based on these we moved on to calculate the density and displacement maps of the SIR3 molecules. Here it should be noted that the tracking of SIR3 is performed in 2-dimensions and the molecules are observable as long as are inside the focal plan which is the z-section of about 400 nm (Figure 2D). After measuring all the traces, we computed the Probability Density Function for the trace lengths, and here we found that while the shortest traces seemed to follow an exponential decay, there was a tail with some very long traces (Figure 2E). Here it is important to note that the half-life time of JF6467 is approximately 2 seconds, meaning that the short traces are due to molecules moving out of the observable z-section and not the photo bleaching of the JF647 dyes.

We aimed to estimate the effective diffusion coefficient of SIR3 in the WT environment, and therefore we computed the displacement for all points in each trace separately, and grouped these into the displacement histogram (Hansen et al., 2018; Klein et al., 2019; Stracy and Kapanidis, 2017). In this way we could test the naive hypothesis that SIR3 molecules simply exhibit a single diffusive motion. We therefore fit the displacement histograms to a Rayleigh distribution (a one parameter fit), and use the resulting fit quality to determine if this hypothesis is sufficient to describe the obtained data. By using Maximum-likelihood minimisation, we extract the most likely value for the diffusion coefficient and based on this we use the Kolmogorov-Smirnoff (KS) test, obtaining a p-value of  $p = 0.0001$ , indicating that more complex motion takes place. To take into account that the molecules can diffuse inside the silencing foci and outside these, we introduce two subpopulations characterized by distinct diffusion coefficients (see Materials and methods). By analysing individual cells, we observe that single traces can be very long in a small region of space, indicating a lower diffusion coefficient (Figure 2F). By fitting the displacement histogram with the two-population fit, we reveal that this leads to a good agreement. We further introduce a third population of diffusion coefficients, but obtain similar quality of the fit arguing that two diffusion coefficients is sufficient to explain the motion of the data (Figure 2G).



**Figure 2.** Mobility of Sir3 inside foci in WT cells.

Based on this we conclude that SIR3 in WT seem to have a motion defined by two distinct populations, significantly different from each other. While one of these populations has a very small diffusion coefficient, representing the motion inside the focus, the other seem to be slow compared to free molecules (compare to the free RAD52 for instance in Miné-Hattab et al., 2022). Therefore we hypothesise that this could be related to motion of SIR3 molecules attached to single telomeres, that are not part of the foci and therefore has higher mobility. In order to test this hypothesis, we tried to remove the existing foci and obtain the motion in this environment.

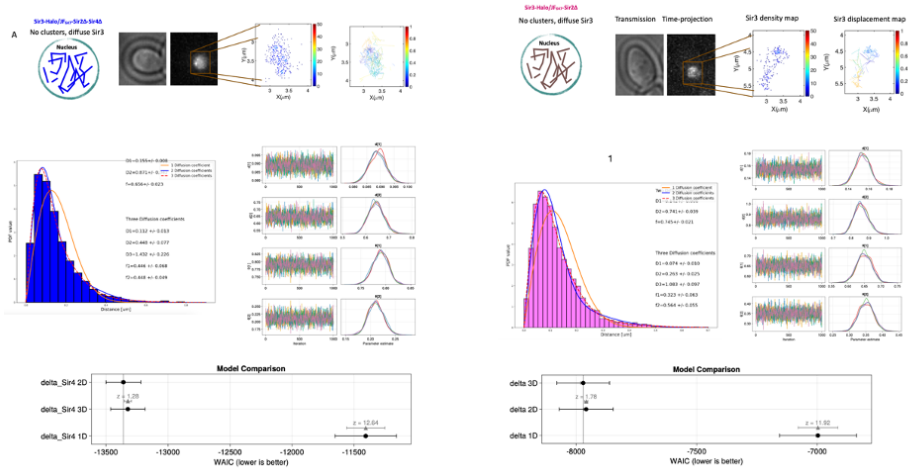
### 168 3.2 | Increased mobility of SIR3 in SIR2D-4D mutants

From the theory of silencing foci, it is well established that the proteins SIR2D and SIR4D should be present in order for the foci to assemble. Therefore, we hypothesised that by deleting these two related genes and thereby removing the availability of SIR2D and SIR4D, the silencing foci should not be able to form (Figure 3A). We succeeded in doing this, and observed that the motion of SIR3

seemed less dense at specific locations compared to the motion of WT (Figure 3B, compare to  
 174 Figure 2B). Furthermore, we also noted that the traces seemed to be shorter and no traces were  
 as long as the ones we observed in the WT conditions (Figure 3B). Therefore again computed the  
 176 displacement histogram and used similar methods as described in Figure 2G to extract the dif-  
 fusion coefficients. Again, we found that one diffusion coefficient could not explain the motion  
 178 of SIR3, but two- and three subpopulations did indeed fit the data sufficiently well. Even though  
 the three-diffusion coefficient fit did lead to a slight improvement in the fit, the two-population  
 180 fitted the data very well (Figure 3C). This was further confirmed by turning to the Bayesian analysis,  
 where we obtain a well-defined and unimodal distributions for each of the fitting parameters in  
 182 the two-population fit (Figure 3D). By comparing the related WAIC scores, we also found that the  
 three-population fit leads to a 1.28  $\sigma$  increase in the fit quality, but since this is within statistical  
 184 uncertainty, we conclude that the two-population fit has the most explanatory power of the ob-  
 served data. By inspecting the diffusion coefficients here we note a very interesting aspect: While  
 186 the slow diffusion coefficient, found in the WT motion, has disappeared in the SIR2D-4D mutant,  
 the high diffusion coefficient for the WT is also identified in the motion of SIR3 in the SIR2D-4D  
 188 mutant, but that a new faster population also has emerged. This supports our hypothesis that  
 the slow observed diffusion coefficient in the WT is a result of the motion inside the foci, but that  
 190 the fast diffusion coefficient does not represent freely diffusing molecules, but rather molecules  
 attached the the semi-mobile telomeres. This also means that effectively all SIR3 molecules are  
 192 bound in the WT suggesting a high number of binding sites and a high binding rate of these sites.

To further support these claims, we constructed a SIR2D mutant, that was deprived of SIR2 but  
 194 still had SIR4 (Figure 3E). Here we again found a well distributed map of SIR 3 (Figure 3F), and by com-  
 puting the displacement histogram we revealed that approximately the same diffusion coefficients  
 196 existed in this mutant (Figure 3G – compare to Figure 3C). Here is seemed that the two-population  
 fit differed slightly more from the three-population fit than the double mutant. To compare the  
 198 quality of all hypotheses we again turned to the Bayesian analysis, where we could again find  
 that all parameters in the two-population fit where smoothly, unmorally distributed, and while the  
 200 three population fit had slightly better predictive power it was still only 1.78  $\sigma$  and therefore within  
 statistical uncertainty (Figure 3H).

202 Based on this, we conclude that by depriving the cell of SIR2 (and SIR4), the foci disappears and  
 the mobility of SIR3 is increased and is described by two populations: A slow population repre-

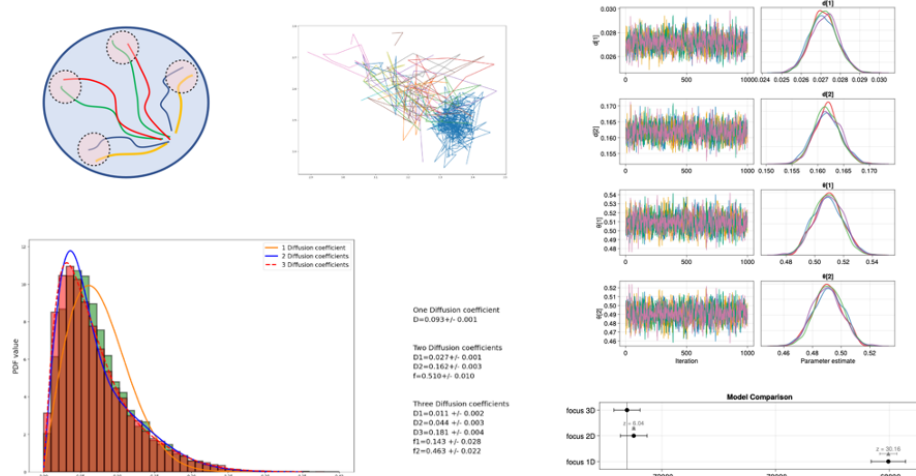


**Figure 3.** Mobility of Sir3 inside foci in mutant strains.

senting the bound molecules to single telomeres and a fast population representing the free SIR3 molecules. To understand the nature of the actual foci, we needed to understand the mobility of SIR3 inside the foci better. Therefore, we investigated the movement of the foci itself.

### 3.3 | Mobility of silencing foci is comparable to the motion of SIR3 inside the foci

Our aim was now to extract the motion of the foci as a single structure and compare this to the motion of the single molecules. In order to obtain this, we used high photo-activation illumination to simultaneously activate all SIR3-mMaple and image the silencing foci as a single entities. Here we are aware that the observed movement should now be dominated by the focus, but since the binding of single SIR3 molecules to the single telomeres, we should be aware that this could also be observed in the data (Figure 4A). We extracted the traces of these whole-mobility structures, and we obtained some confined slowly diffusing traces (blue part in Figure 4B) but also many faster moving traces (multiple colours in Figure 4B). By eye, this does suggest that some movement takes place as a well-defined structure (a silencing-focus) while other motion might be due to the more mobile single telomeres. To test this, we now generated the displacement histograms for the entities, and extracted the diffusion coefficients (Figure 4C). In order to compare the hypotheses of the subpopulations, we directly applied the bayesian analysis and while the two-parameter fit did again lead to well-defined parameters, the three-population fit did lead to a better fit ( $6.08 \sigma$ ). We



**Figure 4.** Individual Sir3 vs. whole focus.

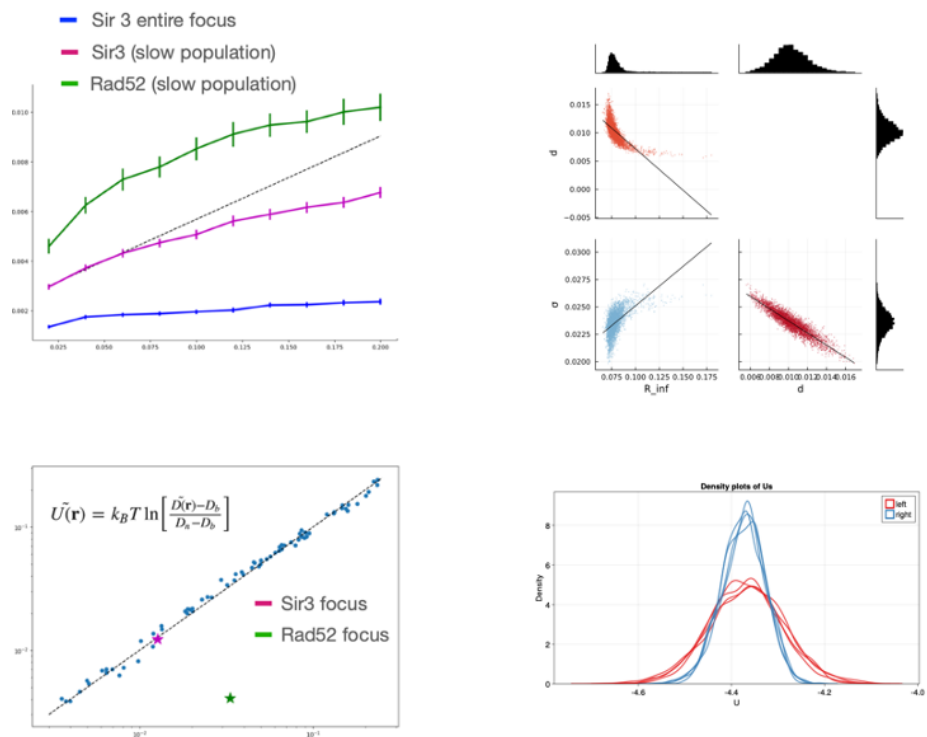
note however that in both fits, the fast part of the population does match the diffusion coefficient of the single telomeres observed for single molecule SIR3 in both WT and SIR2D-SIR4D mutants. Now focusing on the slow part of the diffusion coefficients, we note that these match the extracted diffusion coefficients we found for the single molecule movement of SIR3. However to obtain a better understanding for the similarities of these, and in particular in order to extract the experimental noise levels in the measurements since these might differ significantly for the measurements of the entire focus and and measurements of the single SIR3, we moved on to measure the mean squared distance (MSD) and use these to extract the actual diffusion coefficients.

### 3.4 | Diffusion of SIR3 inside the silencing focus match the predicted movement of a Polymer Bridging Model

Our aim was now to extract the motion of the foci as a single structure and compare this to the motion of the single molecules. In order to obtain this, we used the previously derived theoretical result that connects the diffusion coefficient inside the foci structures to the free energy of these (Heltberg et al., 2021). Here the exact diffusion coefficient is extremely important and the result we obtained in section two is affected by the experimental noise level and this has a significant impact since the diffusion coefficient is so low. In order to separate these we used the method of Mean-Square Distances (MSD). Here we take the slow part of the population in the WT data, and for traces belonging to this family of diffusion coefficients we generate the mean square distances. Finally,

we fit the three first datapoints to a straight line, and use the slope as the diffusion coefficient whereas the intersection is a parameter determined by the experimental noise level. In order to obtain the free energy, we compare the fraction of traces belonging to the slow population, relative to the fast part of the population (See Miné-Hattab et al., 2022 for similar application). In this we take the size of the observable frame compared to the overall size of the cell nucleus into account, as well as we estimate an average of four foci on average. With this we obtain a relation between the free energy and the diffusion coefficient. We know that in a polymer bridging model this should scale as:

$$U = k_{\text{BT}} \ln \left( \frac{D_{\text{inside}} - D_{\text{focus}}}{D_{\text{outside}} - D_{\text{focus}}} \right). \quad (7)$$



**Figure 5.** Free energy and diffusion relation. Relation to Rad52.

232 Here we assume that the diffusion coefficient of the focus is similar to the diffusion coefficient  
233 of the binding sites that would diffuse in a bridging model. We then used the simulation results of  
234 Heltberg et al., 2021, to show that in the simulations this type of structure always yield this relation  
and we showed the result of the relation in the repair foci that is markedly off this line (Figure 5B).

<sup>236</sup> Finally we plotted the result for the silencing foci for the values obtained in this study and here  
<sup>238</sup> we obtained a remarkable agreement. To further validate these results we used the Bayesian  
<sup>240</sup> approach, where we tested the mutual correlation of the parameters investigated in the MSD curve  
<sup>242</sup> (Figure 5C). Next we used this method to extract the free energy from the populations and compare  
<sup>244</sup> this to the free energy estimate based on the extracted diffusion coefficients (Figure 5D). These  
<sup>246</sup> were completely comparable, which further strengthens the conclusion that the motion inside the  
<sup>248</sup> silencing foci is really comparable to what would be theoretically expected in the polymer bridging  
<sup>250</sup> model.

## <sup>244</sup> 4 | DISCUSSION

The two leading hypotheses for describing the nature of nuclear foci is the polymer bridging model  
<sup>246</sup> and the liquid droplet model. In this work we have used the data obtained from SPT experiments  
<sup>248</sup> to investigate the underlying nature of the silencing foci, experienced by the motion of SIR3. We  
<sup>250</sup> find that the behaviour is comparable with the theoretical expectations of the polymer bridging  
<sup>252</sup> model and this work therefore strengthens the hypothesis that these structures are indeed a dense  
<sup>254</sup> collection of binding sites.

From a theoretical perspective, it is noteworthy that the method we apply here cannot directly  
<sup>256</sup> falsify the hypothesis of a liquid structure, but rather it fails to disprove the hypothesis of a polymer  
<sup>258</sup> bridging structure. We use a statistical mechanics formulation, derived a mean field, for the PBM.  
<sup>260</sup> This shares the same functional form as the LPM in the sense that the diffusion coefficient follow a  
<sup>262</sup> step function with one value inside the focus and another value outside. However for the PBM we  
<sup>264</sup> have an additional constraint that precisely links the concentration of proteins and their relative  
<sup>266</sup> diffusion inside the focus: the more time spent inside the focus, the slower the effective diffusion.  
<sup>268</sup> In this sense, if the diffusion coefficient is higher than some value (To the right of the diagonal in  
<sup>270</sup> Figure 4A), then this would typically represent a liquid droplet where diffusion can be faster.  
<sup>272</sup> From a functional perspective, it is also interesting to consider the role of a polymer bridging  
<sup>274</sup> model, compared to a liquid model. The simplest form of a silencing foci, would simply keep away  
<sup>276</sup> the transcription factors and activators. We have previously shown (Heltberg et al., 2021) that the  
<sup>278</sup> existence of a polymer bridging model, would typically increase the first passage time to find a  
<sup>280</sup> target, whereas a liquid model could greatly enhance this. Therefore it is a tempting hypothesis,

that foci formed with the aim of slowing down rates would be of a polymer bridging model, whereas  
266 the foci with the aim of increasing rates (for instance in the repair foci) could be liquid droplets. On a  
more general point, foci are formed inside the nucleus for various reasons with different roles, and  
268 it is clear that they can remain stable very different timescales. Here it is interesting that repair foci  
are maintained for relative short periods (timescale of hours) and they have the ability to quickly  
270 dissolve as long-term stability is not so important. On the other hand, gene expression foci can be  
very stable (Hnisz et al., 2017; Bing et al., 2020), and this could be explained by the hypothesis that  
272 these would typically be polymer bridging structures.

#### 4.1 | Acknowledgment

274 Acknowledgements here

#### 4.2 | Data availability

276 Source code is hosted at GitHub: <https://github.com/ChristianMichelsen/diffusion>.

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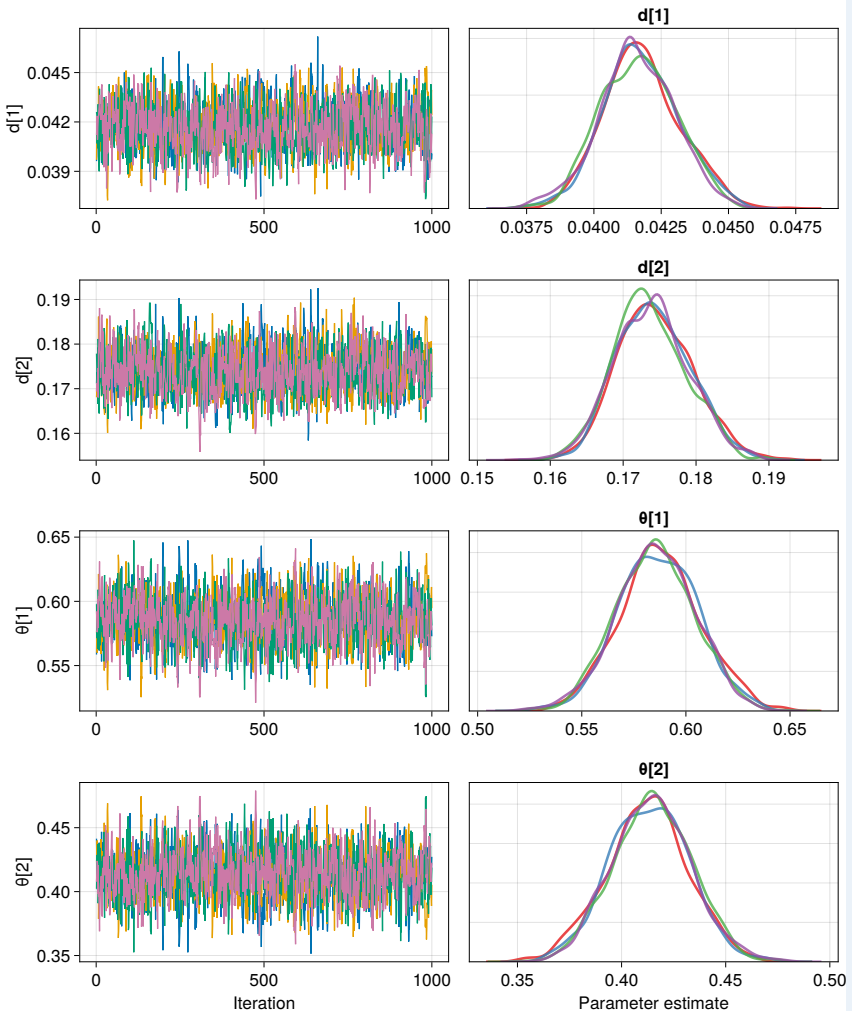
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- <sup>378</sup>

382 Appendix 1

## A | APPENDIX FIGURE 1

384 Here an example of an appendix figure.



386 Appendix 1—figure 1. XXX.



## APPENDIX



## A *Kap København*

The following pages contain the paper published in Nature 2022:

Kurt H. Kjær, Mikkel W. Pedersen, Bianca De Sanctis, Binia De Cahsan, Thorfinn S. Korneliussen, **Christian Michelsen**, Karina K. Sand, Stanislav Jelavić, Anthony H. Ruter, Astrid M. Z. Bonde, Kristian K. Kjeldsen, Alexey S. Tesakov, Ian Snowball, John C. Gosse, Inger G. Alsos, Yucheng Wang, Christoph Dockter, Magnus Rasmussen, Morten E. Jørgensen, Birgitte Skadhauge, Ana Prohaska, Jeppe Å. Kristensen, Morten Bjerager, Morten E. Allentoft, Eric Coissac, PhyloNorway Consortium, Alexandra Rouillard, Alexandra Simakova, Antonio Fernandez-Guerra, Chris Bowler, Marc Macias-Fauria, Lasse Vinner, John J. Welch, Alan J. Hidy, Martin Sikora, Matthew J. Collins, Richard Durbin, Nicolaj K. Larsen & Eske Willerslev, “*A 2-million-year-old ecosystem in Greenland uncovered by environmental DNA*” (Published in Nature, 2022, doi: 10.1038/s41586-022-05453-y).

The paper use the metaDMG tool to identify ancient species and classify the amount of ancient damage in these species. This shows, that modern modern statistical methods combined with excellent work in the ancient DNA labs can provide new insights into the past – even on more than two millions years old data.

## Article

# A 2-million-year-old ecosystem in Greenland uncovered by environmental DNA

<https://doi.org/10.1038/s41586-022-05453-y>

Received: 30 September 2021

Accepted: 18 October 2022

Open access

 Check for updates

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Late Pliocene and Early Pleistocene epochs 3.6 to 0.8 million years ago<sup>1</sup> had climates resembling those forecasted under future warming<sup>2</sup>. Palaeoclimatic records show strong polar amplification with mean annual temperatures of 11–19 °C above contemporary values<sup>3,4</sup>. The biological communities inhabiting the Arctic during this time remain poorly known because fossils are rare<sup>5</sup>. Here we report an ancient environmental DNA<sup>6</sup> (eDNA) record describing the rich plant and animal assemblages of the Kap København Formation in North Greenland, dated to around two million years ago. The record shows an open boreal forest ecosystem with mixed vegetation of poplar, birch and thuja trees, as well as a variety of Arctic and boreal shrubs and herbs, many of which had not previously been detected at the site from macrofossil and pollen records. The DNA record confirms the presence of hare and mitochondrial DNA from animals including mastodons, reindeer, rodents and geese, all ancestral to their present-day and late Pleistocene relatives. The presence of marine species including horseshoe crab and green algae support a warmer climate than today. The reconstructed ecosystem has no modern analogue. The survival of such ancient eDNA probably relates to its binding to mineral surfaces. Our findings open new areas of genetic research, demonstrating that it is possible to track the ecology and evolution of biological communities from two million years ago using ancient eDNA.

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The Kap København Formation is located in Peary Land, North Greenland (82° 24' N 22° 12' W) in what is now a polar desert. The upper depositional sequence contains well-preserved terrestrial animal and plant remains washed into an estuary during a warmer Early Pleistocene interglacial cycle<sup>7</sup> (Fig. 1). Nearly 40 years of palaeoenvironmental and climate research at the site provide a unique perspective into a period when the site was situated at the boreal Arctic ecotone with reconstructed summer and winter average minimum temperatures of 10 °C and -17 °C respectively—more than 10 °C warmer than the present<sup>7–11</sup>.

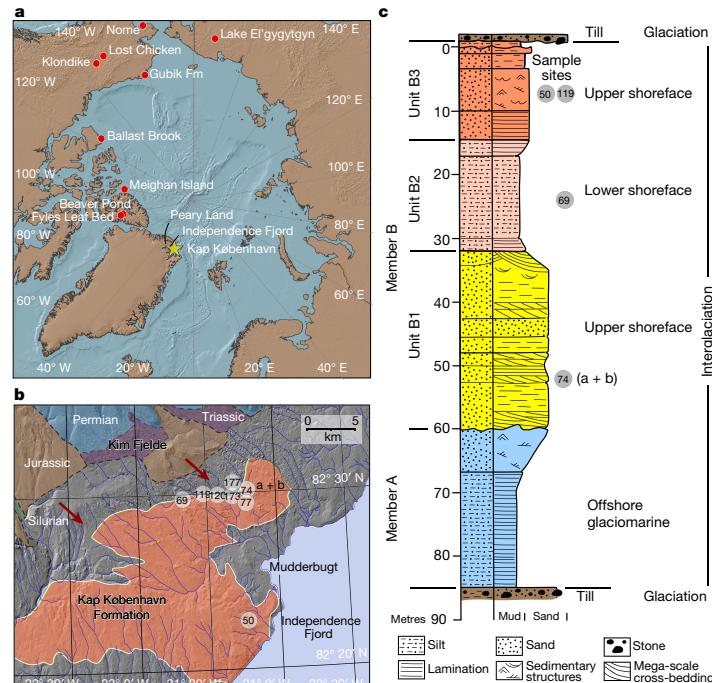
These conditions must have driven substantial ablation of the Greenland Ice Sheet, possibly producing one of the last ice-free intervals<sup>7</sup> in the last 2.4 million years (Myr). Although the Kap København Formation is known to yield well-preserved macrofossils from a coniferous boreal forest and a rich insect fauna, few traces of vertebrates have been found. To date, these comprise remains from lagomorph genera, their coprolites and *Aphodius* beetles, which live in and on mammalian dung<sup>10,11</sup>. However, the approximately 3.4 Myr old Fyles Leaf bed and Beaver Pond on Ellesmere Island in Arctic Canada preserve fossils of

[Q5]

[Q6]

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## Article



**Fig. 1 | Geographic location and depositional sequence.** **a**, Location of Kap København Formation in North Greenland at the entrance to the Independence Fjord ( $82^{\circ}24'N$ ,  $22^{\circ}12'W$ ) and locations of other Arctic Plio-Pleistocene fossil-bearing sites (red dots). **b**, Spatial distribution of the erosional remnants of the 100-m thick succession of shallow marine near-shore sediments between Mudderbugt and the low mountains towards the north. **c**, Glacial-interglacial

mammals that potentially could have colonized Greenland, such as the extinct bear (*Protartos abstrusus*), giant beavers (*Dipoides* sp.), the small canine *Eucyon* and Arctic giant camelines<sup>4,12,13</sup> (similar to *Paracamelus*). Whether the Nares Strait was a sufficient barrier to isolate northern Greenland from colonization by this fauna remains an open question.

The Kap København Formation is formally subdivided into two members<sup>7</sup> (Fig. 1). The lower Member A consists of up to 50 m of laminated mud with an Arctic ostracod, foraminifera and mollusc fauna deposited in an offshore glaciomarine environment<sup>14</sup>. The overlying Member B consists of 40–50 m of sandy (units B1 and B3) and silty (unit B2) deposits, including thin organic-rich beds with an interglacial macrofossil fauna that were deposited closer to the shore in a shallow marine or estuarine environment represented by upper and lower shoreface sedimentary facies<sup>7</sup>.

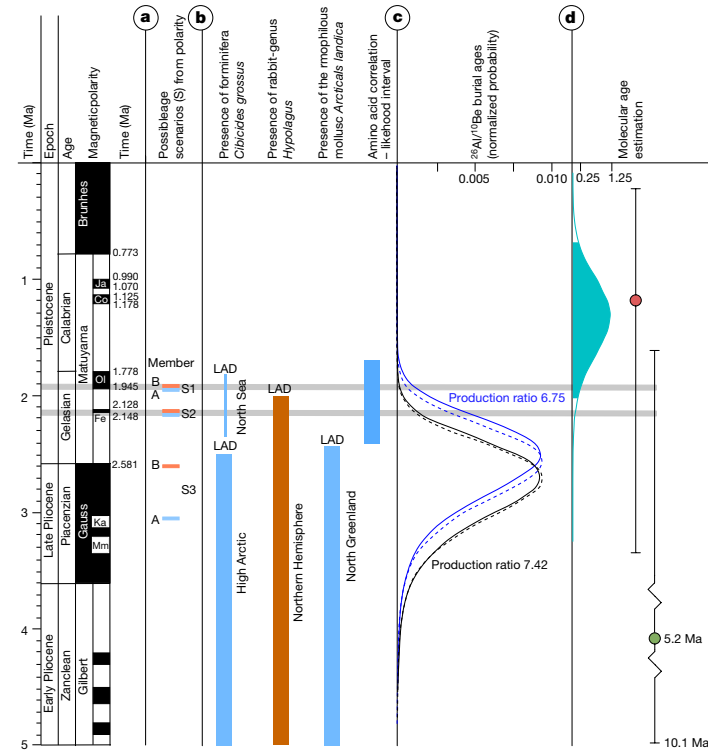
The specific depositional environments are also reflected in the mineralogy of the units, where the proximal B3 locality has the lowest clay and highest quartz contents (Sample compositions in Supplementary Tables 4.2.1 and 4.2.2 and unit averages in Supplementary Tables 4.2.3 and 4.2.4). The architecture of the basin infill suggests that Member B units thicken towards the present coast—that is, distal to the sediment source in the low mountains in the north (Fig. 1). Abundant organic detritus horizons are recorded in units B1 and B3, which also contain beds rich in arctic and boreal plant and invertebrate macrofossils, as well as terrestrial mosses<sup>10,15</sup>. Therefore, the taphonomy of the DNA

division of the depositional succession of clay Member A and units B1, B2 and B3 constituting sandy Member B. Sampling intervals for all sites are projected onto the sedimentary succession of locality 50. Sedimentological log modified after ref. <sup>7</sup>. Circled numbers on the map mark sample sites for environmental DNA analyses, absolute burial dating and palaeomagnetism. Numbered sites refer to previous publications<sup>7,10,11,14,98</sup>.

most probably reflects the biological communities eroded from a range of habitats, fluvially transported to the foreshore and concentrated as organic detritus mixed into sandy near-shore sediments within units B1 and B3. Conversely, the deeper water facies from Member A and unit B2 have a stronger marine signal. This scenario is supported by the similarities in the mineralogic composition between Kap København Formation sediments and Kim Fjelde sediments (Supplementary Tables 4.2.1 and 4.2.5).

## Geological age

A series of complementary studies has successively narrowed the depositional age bracket of the Kap København Formation from 4.0–0.7 million years ago (Ma) to a 20,000-year-long age bracket around 2.4 Ma (see Supplementary Information, sections 1–3). This was achieved by a combination of palaeomagnetism, biostratigraphy and allostratigraphy<sup>7,14,16–18</sup>. Notably, the last appearance data of the mammals, foraminifera and molluscs in the stratigraphic record show an age close to 2.4 Myr (see Supplementary Information, section 2). Within this overall framework, we add new palaeomagnetic data showing that Member A has reversed magnetic polarity and the main part of the overlying unit B2 has normal magnetic polarity. In the context of previous work, this is consistent with three magnetostratigraphic intervals in the Early Pleistocene where there is a reversal: 1.93 Myr (scenario 1), 2.14 Myr (scenario 2) or 2.58 Myr (scenario 3) (Supplementary



**Fig. 2 | Age proxies for the Kap København Formation.** **a**, Revised palaeomagnetic analysis shows unit B2 to have normal polarity and unlocks three possible age scenarios (S1–S3) including Members A (blue) and B (brown). Normal polarity is coloured black and reverse polarity is shown in white. Ja, Jaramillo; Co, Cobb Mountain; Ol, Olduvai; Fe, Feni; Ka, Kaena; Mm, Mammoth. **b**, Presence and last appearance datum (LAD) for marine foraminifera *Cibicides grossus*, rabbit-genus *Hypolagus* and the mollusc *Arctica islandica* in the High Arctic, Northern Hemisphere and North Greenland, respectively. The blue band on the far right indicates the age range for Member A estimated from amino acid ratios on shells<sup>7</sup>. **c**, Convolved probability distribution functions for cosmogenic burial ages calculated for two different production ratios

(7.42 (black) and 6.75 (blue)). The dashed line and the solid line show the distributions for steady erosion and zero erosion, respectively. These distributions are all maximum ages. **d**, Molecular dating of *Betula* sp., yielding a median age of the DNA in the sediment of 1.323 Myr, with whiskers confining the 95% height posterior density (HPD) of 0.68 to 2.02 Myr (blue density plot), running Markov chain Monte Carlo estimation over for 100 million iterations. The red dot is the median molecular age estimate found using the Mastodon mitochondrial genome restricting to radiocarbon-dated specimens, whereas the green area includes molecular clock estimated specimens in BEAST, running Markov chain Monte Carlo estimation for 400 million iterations. Whiskers confine the 95% HPD.

Information, section 1). Furthermore, we constrain the age using cosmogenic  $^{26}\text{Al}$ : $^{10}\text{Be}$  burial dating of Member B at four sites in this study (Supplementary Information, section 3). The recommended maximum burial age for the Kap København Formation is  $2.70 \pm 0.46$  Myr (Fig. 2; Methods). However, we discard the older scenario 3 as it contradicts the evidence for a continuous sedimentation across Members A and B during a single glacial–interglacial depositional cycle<sup>7,14,16,18,19</sup>. This leaves two possible scenarios (scenarios 1 and 2), in which scenario 1 supports an age of 1.9 Myr and scenario 2 supports an age of 2.1 Myr.

### DNA preservation

DNA degrades with time owing to microbial enzymatic activity, mechanical shearing and spontaneous chemical reactions such as hydrolysis and oxidation<sup>20</sup>. The oldest known DNA obtained to date has been recovered from a permafrost-preserved mammoth molar dated to 1.2–1.1 Ma using geological methods and 1.7 Ma (95% highest posterior density, 2.1–1.3 Ma) using molecular clock dating<sup>21</sup>. To explore the

likelihood of recovering DNA from sediments at the Kap København formation, we calculated the thermal age of the DNA and its expected degree of depurination at the Kap København Formation. Using the mean average temperature<sup>22</sup> (MAT) of  $-17^\circ\text{C}$ , we found a thermal age of  $2.7 \text{ kyr}_{\text{DNA}@10^\circ\text{C}}$ —that is, 741 times less than the age of 2.0 Myr (Supplementary Information, section 4 and Supplementary Table 4.4.1). Using the rate of depurination from Moa bird fossils<sup>23</sup>, we found it plausible that DNA with an average size of 50 base pairs (bp) could survive at the Kap København Formation, assuming that the site remained frozen (Supplementary Information, section 4 and Supplementary Table 4.4.2). Mechanisms that preserve DNA in sediments are likely to be different from that of bone. Adsorption at mineral surfaces modifies the DNA conformation, probably impeding molecular recognition by enzymes, which effectively hinders enzymatic degradation<sup>24–27</sup>. To investigate whether the minerals found in Kap København Formation could have retained DNA during the deposition and preserved it, we determined the mineralogic composition of the sediments using X-ray diffraction and measured their adsorption capacities. Our findings

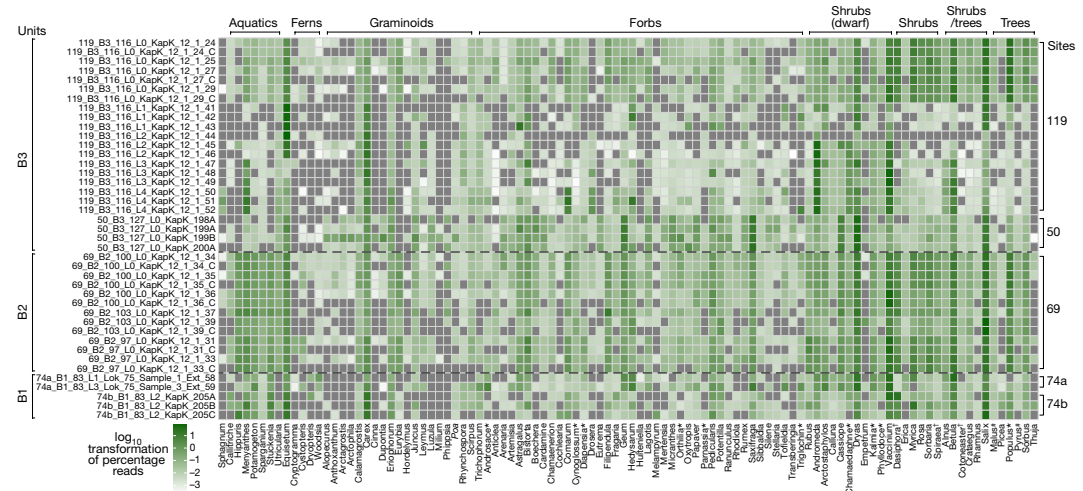
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## Article



**Fig. 3 | Early Pleistocene plants of Northern Greenland.** Metagenomic taxonomic profiles of the plant assemblage. Taxa in bold are genera only found as DNA and not as macrofossil or pollen. Asterisks indicate those that are found at other Pliocene arctic sites. Extinct species as identified by either

macrofossils or phylogenetic placements are marked with a dagger. Reads classified as *Pyrus* and *Malus* are marked with a pound symbol, and are probably over-classified DNA sequences belonging to another species within Rosaceae that are not present as a reference genome.

highlight that the marine depositional environment favours adsorption of extracellular DNA on the mineral surfaces (Supplementary Information, section 4 and Supplementary Table 4.3.1.1). Specifically, the clay minerals (9.6–5.5 wt%) and particularly smectite (1.2–3.7 wt%), have higher adsorption capacity compared to the non-clay minerals (59–75 wt%). At a DNA concentration representative of the natural environments<sup>28</sup> (4.9 ng ml<sup>-1</sup> DNA), the DNA adsorption capacity of smectite is 200 times greater than for quartz. We applied a sedimentary eDNA extraction protocol<sup>29</sup> on our mineral-adsorbed DNA samples, and retrieved only 5% of the adsorbed DNA from smectite and around 10% from the other clay minerals (Methods and Supplementary Information, section 4). By contrast, we retrieved around 40% of the DNA adsorbed to quartz. The difference in adsorption capacity and extraction yield from the different minerals demonstrates that mineral composition may have an important role in ancient eDNA preservation and retrieval.

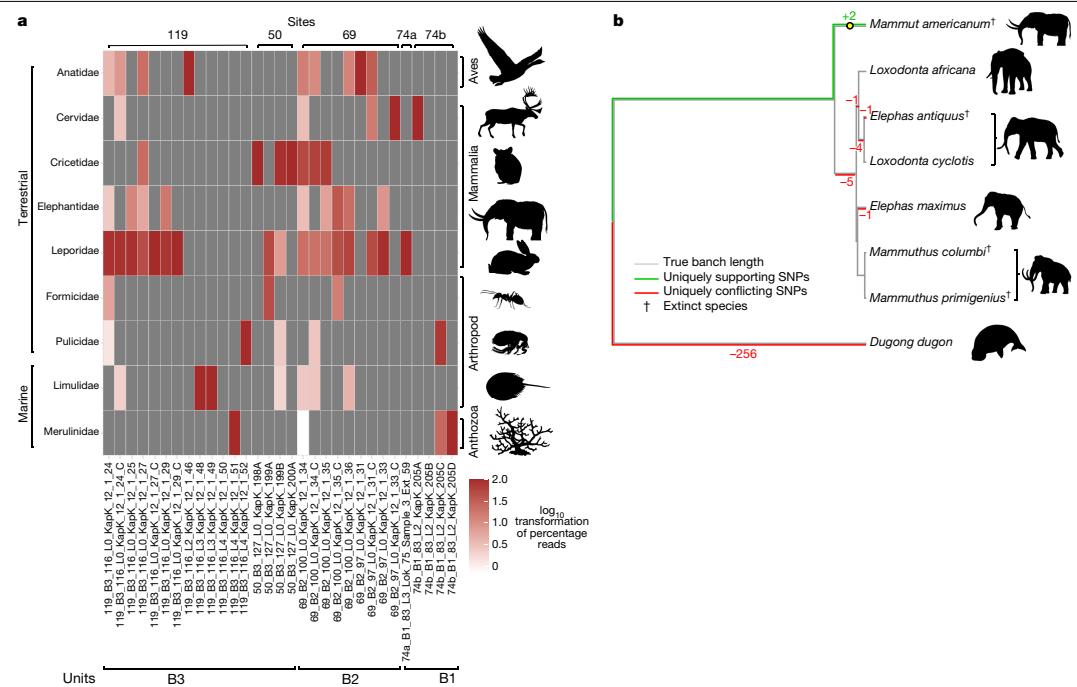
### Kap København metagenomes

We extracted DNA<sup>29</sup> from 41 organic-rich sediment samples at five different sites within the Kap København Formation (Supplementary Information, section 6 and Source Data 1), which were converted into 65 dual-indexed Illumina sequencing libraries<sup>30</sup>. First, we tested 34 of the 65 libraries for plant plastid DNA by screening for the conserved photosystem II D2 (*psbD*) gene using droplet digital PCR (ddPCR) with a gene-targeting primer and probe spanning a 39-bp region and a P7 index primer. Further, we screened for the *psbA* gene using a similar assay targeting the Poaceae (Methods and Supplementary Fig. 6.12.1). A clear signal in 31 out of 34 samples tested confirmed the presence of plant plastid DNA in these libraries (Source Data 1, sheets 5 and 6). Additionally, we subjected 34 of the 65 libraries to mammalian mtDNA capture enrichment using the Arctic PaleoChip 1.0<sup>31</sup> and shotgun sequenced all libraries (initial and captured) using the Illumina HiSeq 4000 and NovaSeq 6000. A total of 16,882,114,068 reads were sequenced, which after adaptor trimming, filtering for ≥30 bp and a minimum phred quality of 30 and duplicate removal resulted in 2,873,998,429 reads. These

were analysed for kmer comparisons using simka<sup>32</sup> (Supplementary Information, section 6) and then parsed for taxonomic classification using competitive mapping with HOLI (<https://github.com/miwipe/KapCopenhagen.git>), which includes a recently published dataset of more than 1,500 genome skims of Arctic and boreal plant taxa<sup>33,34</sup> (Methods and Supplementary Information, section 6). Considering the age of the samples and thus the potential genetic distance to recent reference genomes, we allowed each read to have a similarity between 95–100% for it to be taxonomically classified using ngsLCA<sup>35</sup>. The metaDMG (v.0.14.0) program (<https://metadmg-dev.github.io/metaDMG-core/index.html>) was subsequently used to quantify and filter each taxonomic node for postmortem DNA damage for all the metagenomic samples (Methods). This method estimates the average damage at the termini position (D-max) and a likelihood ratio (λ-LR) that quantifies how much better the damage model (that is, more damage at the beginning of the read) fits the data compared with a null model (that is, a constant amount of damage; see Supplementary Information, section 6). We found the DNA damage to be highly increased, especially for eukaryotes (mean D-max = 40.7%). From this we set D-max ≥25% as a filtering threshold for a taxonomic node to be parsed for further downstream analysis as well as a λ-LR higher or equal to 1.5. We furthermore set a threshold requiring that the minimum number of reads per taxon exceeded the median of reads assigned across all taxa divided by two to filter for taxa in low abundance. Similarly, for a sample to be considered, the total number of reads for a sample had to exceed the median number of reads per sample divided by two, to filter for samples with fewest reads. Lastly, we filtered out taxa with fewer than three replicates and subsequently reads were normalized by conversion to proportions (Figs. 3 and 4a).

### DNA, pollen and macrofossils comparison

Greenland's coasts extend from around 60° to 83° N and include bioclimatic zones from the subarctic to the northern polar desert<sup>36,37</sup>. There are 175 vascular plant genera native to Greenland, excluding historically introduced species<sup>38–40</sup>. Of these, 70 (40%) were detected



**Fig. 4 | Early Pleistocene animals of Northern Greenland. a,** Metagenomic taxonomic profiles of the animal assemblage from units B1, B2 and B3. Taxa in bold are genera only found as DNA. **b,** phylogenetic placement and pathPhynder<sup>88</sup> results of mitochondrial reads uniquely classified to Elephantidae or lower (Source Data 1).

by the metagenomic analysis (Fig. 3); the majority of these genera are today confined to bioclimatic zones well to the south of Kap København's polar desert (see ref.<sup>41</sup> and references therein), for example, all aquatic macrophytes. Reads assigned to *Salix*, *Dryas*, *Vaccinium*, *Betula*, *Carex* and *Equisetum* dominate the assemblage, and of these genera, *Equisetum*, *Dryas*, *Salix arctica* and two species of *Carex* (*Carex nardina* and *Carex stans*) grow there currently, whereas only a few records of *Vaccinium uliginosum* are found above 80° N, and *Betula nana* are found above 74° N (ref.<sup>42</sup>). Out of the 102 genera detected in the Kap København ancient eDNA assemblage, 39% no longer grow in Greenland but do occur in the North American boreal (for example, *Picea* and *Populus*) and northern deciduous and maritime forests (for example, *Crataegus*, *Taxus*, *Thuya* and *Filipendula*). Many of the plant genera in this diverse assemblage do not occur on permafrost substrates and require higher temperatures than those at any latitude on Greenland today.

In addition to the DNA, we counted pollen in six samples from locality 119, unit B3 (Methods and Supplementary Fig. 4.1.1). Percentages were calculated for 4 of the samples with pollen sums ranging from 71–225 terrestrial grains (mean = 170.25). Upland herbs, including taxa in the Cyperaceae, Ericales and Rosaceae comprised around 40% of sample 4. Samples 5 and 6 were dominated by arboreal taxa, particularly *Betula*. The Polypodiopsida (for example, *Equisetum*, *Asplenium* and *Athyrium filix-femina*) and Lycopodiopsida (*Lycopodium annotinum* and *Selaginella rupestris*) were also well represented and comprised over 30% of the assemblage in samples 1, 4 and 6.

A total of 39 plant genera out of the 102 identified by DNA also occurred as macrofossils or pollen at the genus level. A further 39 taxa were potentially identified as macrofossil or pollen but not to the same

taxonomic level<sup>10,15</sup> (Source Data 1, sheets 1 and 2). For example, 12 genera of Poaceae were identified by DNA (*Alopecurus*, *Anthoxanthum*, *Arctagrostis*, *Arctophila*, *Calamagrostis*, *Cinna*, *Dupontia*, *Hordelymus*, *Leymus*, *Milium*, *Phippsia* and *Poa*), of these only *Hordelymus* is not found in the Arctic today (<http://panarcticflora.org/>), but these were only distinguished to family level in the pollen analysis and only one Poaceae macrofossil was found. There were 24 taxa that were recorded only as DNA. These included the boreal tree *Populus* and a few shrubs and dwarf shrubs, but mainly herbaceous plants. Of the 73 plant genera recovered as macrofossils<sup>10,15</sup>, only 24 were not detected in the DNA analysis. Because macrofossils and DNA have similar taphonomies—as both are deposited locally—more overlap is expected between them than between DNA and pollen, which is typically dispersed regionally<sup>43</sup>. Nine of the taxa absent in DNA were bryophytes, probably owing to poor representation of this group within the genomic reference databases. Furthermore, the extinct taxon Araceae is not present in the reference databases. The remaining undetected genera were vascular plants, and all except two (*Oxyria* and *Cornus*) were rare in the macrofossil record. Because the detection of rare taxa is challenging in both macrofossil and DNA records<sup>44</sup>, we argue that this overlap between the DNA and macrofossil records is as high as can be expected on the basis of the limitations of both methods.

An additional 19 taxa were recorded in the pollen record presented here and in that of Bennike<sup>45</sup> including four trees or shrubs, five ferns, three club mosses, and one each of algae, fungi and liverwort. We also find pollen from anemophilous trees, particularly gymnosperms, which can be distributed far north of the region where the plants actually grow<sup>10</sup>. Bennike<sup>45</sup> also notes a high proportion of club mosses and ferns and suggests they may be overrepresented owing to their spore wall

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being resistant to degradation. Furthermore, if these taxa were preferentially distributed along streams flowing into the estuary, their spores could be relatively more concentrated in the alluvium than the pollen of more generally distributed taxa. Thus, both decay resistance and alluvial deposition could contribute to the relative frequencies we observe. This same alluvial dynamic might also have contributed to the very large read counts for *Salix*, *Betula*, *Populus*, *Carex* and *Equisetum* in the metagenomic record, implying that neither the proportion of these taxa in the pollen records nor read counts necessarily correlate with their actual abundance in the regional vegetation in terms of biomass or coverage.

Finally, we sought to date the age of the plant DNA by phylogenetic placement of the chloroplast DNA. We examined data for the genera *Betula*, *Populus* and *Salix*, because these had both sufficiently high chloroplast genome coverage (with mean depth  $24.16 \times$ ,  $57.06 \times$  and  $27.04 \times$ , respectively) and sufficient present-day whole chloroplast reference sequences (Methods). Owing to their age and hence potential genetic distance from the modern reference genomes, we lowered the similarity threshold of uniquely classified reads to 90% and merged these by unit to increase coverage. Both *Betula* and *Salix* placed basally to most of the represented species in the respective genera, and the *Populus* placement results showed support for a mixture of different species related to *P. trichocarpa* and *P. balsamifera* (Extended Data Figs. 7–9).

We used the *Betula* chloroplast reads for a molecular dating analysis, because they were placed confidently on a single edge of the phylogenetic tree (that is, not a mixture as in *Populus*), had a large number of reference sequences, and had high coverage in the ancient sample. We used BEAST<sup>46</sup> v1.10.4 to obtain a molecular clock date estimate for our ancient *Betula* chloroplast sample (see Methods, ‘Molecular dating methods’ for details). We included 31 modern *Betula* and one *Alnus* chloroplast reference sequences, used only sites that had a depth of at least 20 in the ancient sample, and included a previously estimated *Betula*–*Alnus* chloroplast divergence time<sup>47</sup> of 61.1 Myr for calibration of the root node. Our BEAST analysis was robust to both different priors on the age of the ancient sample, and to different nucleotide substitution models (Supplementary Fig. 10). This yielded a median age estimate of 1.323 Myr, with a 95% HPD of (0.6786, 2.0172) Myr (Fig. 2).

### Animal DNA results

The metazoan mitochondrial and nuclear DNA record was much less diverse than that of the plants but contained one extinct family, one that is absent from Greenland today, and four vertebrate genera native to Greenland as well as representatives of four invertebrate families (Fig. 4a). Assignments were based on incomplete and variable representation of reference genomes, so we identified reads to family level, and only where sufficient mitochondrial reads were present, we refined the assignment to genus level by matching these into mitochondrial phylogenies based on more complete present-day mitochondrial sequences (Supplementary Information, section 6). As for the plant reads, uniquely classified animal reads with more than 90% similarity were parsed and merged by unit to increase coverage for phylogenetic placement.

Most notably, we found reads in unit B2 and B3 assigned to the family Elephantidae, which includes elephants and mammoths, but taxonomically not mastodon (*Mammuthus* sp.)—which are, however, in the NCBI taxonomy, and therefore our analysis reads classified to Elephantidae or below therefore include *Mammuthus* sp. A consensus genome of our Elephantidae mitochondrial reads falls on the *Mammuthus* sp. branch (Fig. 4b) and is placed basal to all clades of mastodons. However, we note that this placement within the mastodons depends on only two transition single nucleotide polymorphisms (SNPs), with the first one supported by a read depth of three and the second by only one (Extended Data Fig. 4, Methods and Supplementary Information, section 6). Furthermore, we attempted dating the recovered mastodon

mitochondrial genome using BEAST<sup>48</sup>. We implemented two dating approaches, one was based on using radiocarbon-dated specimens alone, while the other used radiocarbon- and molecular-dated mastodons. The first analysis yielded a median age estimate for our mastodon mitogenome of 1.2 Myr (95% HPD: 191,000 yr–3.27 Myr), the second approach resulted in a median age estimate of 5.2 Myr (95% HPD: 1.64–10.1 Myr) (Supplementary Fig. 6.8.5 and Supplementary Information, section 6).

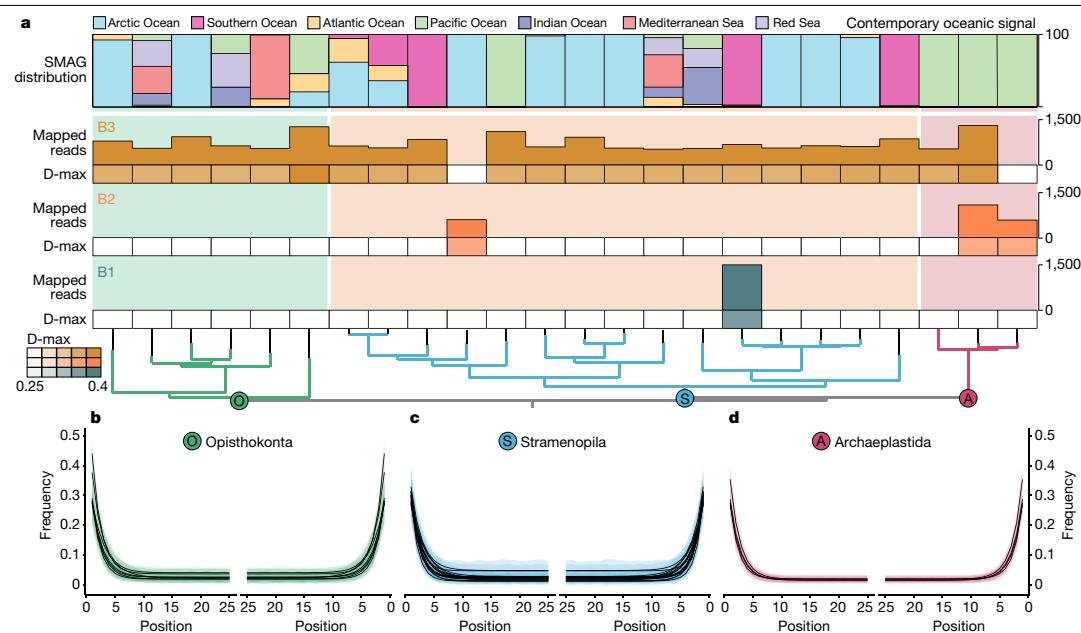
Similarly, reads assigned to the Cervidae support a basal placement on the *Rangifer* (reindeer and caribou) branch (Extended Data Fig. 3). Mitochondrial reads mapping to Leporidae (hares and rabbits) place near the base to the Eurasian hare clade (Extended Data Fig. 2), which is the only mammal found in the fossil record<sup>49</sup>. *Lepus*, specifically *Lepus arcticus*, is also the only genus in the Leporidae living in Greenland today. Mitochondrial reads assigned to Cricetidae cover only one informative transversion SNP, which places them as deriving from the subfamily Arvicolinae (voles, lemmings and muskrats) (Extended Data Fig. 6). For the only avian taxon represented in our dataset—Anatidae, the family of geese and swans—we found a robust basal placement to the genus *Branta* of black geese, supported by three transversion SNPs with read depths ranging between two and four (Extended Data Fig. 5). The refined vertebrate assignments based on mitochondrial references are more biogeographically conserved than for plants. *Dicrostonyx*—specifically *Dicrostonyx groenlandicus* (the Nearctic collared lemming)—is the only genus of the Cricetidae native to Greenland today, just as *Rangifer*—specifically *Rangifer tarandus groenlandicus* (the barren-ground caribou)—is the only member of the Cervidae. The mastodon is the exception, as no member of the Elephantidae lives in present-day Greenland.

### Ancient DNA from marine organisms

The other metazoan taxa identified in the DNA record were a single reef-building coral (Merulinidae) and several arthropods, with matches to two insects—Formicidae (ants) and Pulicidae (fleas)—and one marine family—Limulidae (horseshoe crabs). This is somewhat unexpected, given the rich insect macrofossil record from the Kap København Formation, which comprises more than 200 species, including *Formica* sp. The marine taxa are less abundant than the terrestrial taxa, and no mitochondrial DNA was identified from marine metazoans. The read lengths, DNA damage and the fact that the reads assigned distribute evenly across the reference genomes suggests that these are not artefacts but may be over-matched DNA sequences of closely related, potentially extinct species within the families that are currently absent from our reference databases owing to poor taxonomic representation. By contrast, Limulidae, in the subphylum *Chelicerata*, is unlikely to be misidentified as this distinct genus is the only surviving member within its order and thus deeply diverged from other extant organisms.

The probable source of these reads is a population of *Limulus polyphemus*, the only Atlantic member of the genus, which would have spawned directly onto the sediment as it accumulated. Today this genus does not spawn north of the Bay of Fundy (about 45° N), suggesting warmer surface water conditions in the Early Pleistocene at Kap København consistent with the +8 °C annual sea surface temperature anomaly reconstructed for the Pleistocene of the coast of northeast Greenland<sup>49</sup>. By aligning our reads against the Tara Oceans eukaryotic metagenomic assembled genomes (SMAGs) data (Methods), we further reveal the presence of 24 marine planktonic taxa in 14 samples, covering both zooplankton and phytoplankton (Fig. 5). These detected SMAGs belong to the supergroups Opisthokonta (6), Stramenopila (15) and Archaeplastida (3). The majority of these signals are from SMAGs associated with cold regions in the modern ocean (that is, the Arctic Ocean and Southern Ocean), such as diatoms (Bacillariophyta), Chrysophyceae and the MAST-4 group (Supplementary

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**Fig. 5 | Marine planktonic eukaryotes identified at the Kap København Formation.** **a**, Detection of SMAGs and average damage (D-max) of a SMAG within a member unit. Top, the SMAG distribution in contemporary oceans based on the data of Delmont et al.<sup>73</sup>. The SMAGs are ordered on the basis of phylogenomic inference from Delmont et al.<sup>73</sup>. **b–d**, Distribution of DNA damage among the taxonomic supergroup Opisthokonta (**b**), Stramenopila (**c**) and Archaeplastida (**d**) (Source Data 1).

Table 6.11.1), as we expected. However, a few are cosmopolitan, whereas others, such as Archaeplastida (green microalgae), have an oceanic signal that is today confined to more temperate waters in the Pacific Ocean (Fig. 5). Although we do not know whether modern day ecologies can be extrapolated to ancient ecosystems, the abundance of green microalgae is believed to be increasing in Arctic regions, which tends to be associated with warming surface waters.

## Discussion

The Kap København ancient eDNA record is extraordinary for several reasons; the upper limit of the 95% highest posterior density of the estimated molecular age is 2.0 Myr and independently supports a geological age of approximately 2 Myr (Fig. 2). This implies that the DNA is considerably older than any previously sequenced DNA<sup>21</sup>. Our DNA results detected five times as many plant genera as previous studies using shotgun sequencing of ancient sediments<sup>29,34,50,51</sup>, which is well within the range of the richest northern boreal metabarcoding records<sup>52</sup>. The accuracy of the assignments is strengthened by the observation that 76% of the taxa identified to the level of genus or family also occurred in macrofossil and/or pollen assemblages from the same units. Our results demonstrate the potential of ancient environmental metagenomics to reconstruct ancient environments, phylogenetically place and date ancient lineages from diverse taxa from around 2 Ma (Supplementary Information, section 6). Finally, the DNA identified a set of additional plant genera, which occur as macrofossils at other Arctic Late Pliocene and Early Pleistocene sites (Figs. 1 and 3a and Supplementary Information, section 5) but not as fossils at Kap København, thereby expanding the spatiotemporal distribution of these ancient floras.

Of note, the detection of both *Rangifer* (reindeer and caribou) and *Mammut* (mastodon) forces a revision of earlier palaeoenvironmental reconstructions based on the site's relatively impoverished faunal record, entailing both higher productivity and habitat diversity for much of the deposition period. Because all the vertebrate taxa identified by DNA are herbivores, their representation may be a function of relative biomass (see discussion on taphonomy in Supplementary Information, section 6). Caribou, geese, hares and rodents can all be abundant, at least seasonally, in boreal environments. Additionally, the excrement of large herbivores (such as caribou and particularly mastodons) can be a significant component of sediments<sup>34</sup>. By contrast, carnivores are not represented, consistent with their smaller total biomass. This dynamic also explains the dominance of plant reads over metazoans and to some extent differences in representation of various plant genera (Supplementary Information, section 6). In the general absence of fossils, DNA may prove the most effective tool for reconstructing the biogeography of vertebrates through the Early Pleistocene. DNA from mastodon must imply a viable population of this large browsing megaherbivore, which would require a more productive boreal habitat than that inferred in earlier reconstructions based primarily on plant macrofossils<sup>7</sup>. Mastodon dung from a site in central Nova Scotia from around 75,000 years ago contained macrofossils from sedges, cattail, bulrush, bryophytes and even charophytes, but was dominated by spruce needles and birch samaras<sup>53</sup>. The Kap København units with mastodon DNA yielded macrofossils and DNA from *Betula* as well as more thermophilic arboreal taxa including *Thuja*, *Taxus*, *Cornus* and *Viburnum*, none of which range into Greenland's hydric Arctic tundra or polar deserts today. The co-occurrence of these taxa in multiple units compels a revision of previous temperature estimates as well as the presence of permafrost.

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No single modern plant community or habitat includes the range of taxa represented in many of the macrofossil and DNA samples from Kap København. The community assemblage represents a mixture of modern boreal and Arctic taxa, which has no analogue in modern vegetation<sup>10,15</sup>. To some degree, this is expected, as the ecological amplitudes of modern members of these genera have been modified by evolution<sup>54</sup>. Furthermore, the combination of the High Arctic photoperiod with warmer conditions and lower atmospheric CO<sub>2</sub> concentrations<sup>55</sup> made the Early Pleistocene climate of North Greenland very different from today. The mixed character of the terrestrial assemblage is also reflected in the marine record, where Arctic and more cosmopolitan SMAGs of Ophistokonta and Stramenophila are found together with horseshoe crabs, corals and green microalgae (Archaeplastida), which today inhabit warmer waters at more southern latitudes.

Megaherbivores, particularly mastodons, could have had a significant impact on an interglacial taiga environment, even providing a top-down trophic control on vegetation structure and composition at this high latitude. The presence of mastodons<sup>56,57</sup> coupled with the absence of anthropogenic fire, which has had a role in some Holocene boreal habitats<sup>58</sup>, are important differences. Another important factor is the proximity and biotic richness of the refugia from which pioneer species were able to disperse into North Greenland when conditions became favourable at the beginning of interglacials. The shorter duration of Early Pleistocene glaciations produced less extensive ice sheets allowing colonization from relatively species-rich coniferous-deciduous woodlands in northeastern Canada<sup>12,59</sup>. More extensive glaciation later in the Pleistocene increasingly isolated North Greenland and later re-colonizations were from increasingly distant and/or less diverse refugia.

In summary, we show the power of ancient eDNA to add substantial detail to our knowledge of this unique, ancient open boreal forest community intermixed with Arctic species, a community composition that has no modern analogues and included mastodons and reindeer, among others. Similar detailed flora and vertebrate DNA records may survive at other localities. If recovered, these would advance our understanding of the variability of climate and biotic interactions during the warmer Early Pleistocene epochs across the High Arctic.

### Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41586-022-05453-y>.

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## Article

### Methods

#### Sampling

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Sediment samples were obtained from the Kap København Formation in North Greenland ( $82^{\circ} 24' 00''\text{N}$   $22^{\circ} 12' 00''\text{W}$ ) in the summers of 2006, 2012 and 2016 (see Supplementary Table 3.1.1). Sampled material consisted of organic-rich permafrost and dry permafrost. Prior to sampling, profiles were cleaned to expose fresh material. Samples were hereafter collected vertically from the slope of the hills either using a 10 cm diameter diamond headed drill bit or cutting out  $40 \times 40 \times 40$  cm blocks. Sediments were kept frozen in the field and during transportation to the lab facility in Copenhagen. Disposable gloves and scalpels were used and changed between each sample to avoid cross-contamination. In a controlled laboratory environment, the cores and blocks were further sub-sampled for material taking only the inner part of sediment cores, leaving 1.5–2 cm between the inner core and the surface that provided a subsample of approximately 6–10 g. Subsequently, all samples were stored at temperatures below  $-22^{\circ}\text{C}$ .

We sampled organic-rich sediment by taking samples and biological replicates across the three stratigraphic units B1, B2 and B3, spanning 5 different sites, site: 50 (B3), 69 (B2), 74a (B1), 74b (B1) and 119 (B3). Each biological replicate from each unit at each site was further sampled in different sublayers (numbered L0–L4, Source Data 1, sheet 1).

#### Absolute age dating

In 2014, Be and Al oxide targets from  $8 \times 1$  kg quartz-rich sand samples collected at modern depths ranging from 3 to 21 m below stream cut terraces were analysed by accelerator mass spectrometry and the cosmogenic isotope concentrations interpreted as maximum ages using a simple burial dating approach<sup>1</sup> ( $^{26}\text{Al}:\text{Be}$  versus normalized  $^{10}\text{Be}$ ). The  $^{26}\text{Al}$  and  $^{10}\text{Be}$  isotopes were produced by cosmic ray interactions with exposed quartz in regolith and bedrock surfaces in the mountains above Kap København prior to deposition. We assume that the  $^{26}\text{Al}:\text{Be}$  was uniform and steady for long time periods in the upper few metres of these gradually eroding palaeo-surfaces. Once eroded by streams and hillslope processes, the quartz sand was deposited in sandy braided stream sediment, deltaic distributary systems, or the near-shore environment and remained effectively shielded from cosmic ray nucleons buried (many tens of metres) under sediment, intermittent ice shelf or ice sheet cover, and—at least during interglacials—the marine water column until final emergence. The simple burial dating approach assumes that the sand grains experienced only one burial event. If multiple burial events separated by periods of re-exposure occurred, then the starting  $^{26}\text{Al}:\text{Be}$  before the last burial event would be less than the initial production ratio (6.75 to 7.42, see discussion below) owing to the relatively faster decay of  $^{26}\text{Al}$  during burial, and therefore the calculated burial age would be a maximum limiting age. Multiple burial events can be caused by shielding by thick glacier ice in the source area, or by sediment storage in the catchment prior to final deposition. These shielding events mean that the  $^{26}\text{Al}:\text{Be}$  is lower, and therefore a calculated burial age assuming the initial production ratio would overestimate the final burial duration. We also consider that once buried, the sand grains may have been exposed to secondary cosmogenic muons (their depth would be too great for submarine nucleonic production). As sedimentation rates in these glaciated near-shore environments are relatively rapid, we show that even the muonic production would be negligible (see Supplementary Information). However, once the marine sediments emerged above sea level, in-situ production by both nucleogenic and muogenic production could alter the  $^{26}\text{Al}:\text{Be}$ . The  $^{26}\text{Al}$  versus  $^{10}\text{Be}$  isochron plot reveals this complex burial history (Supplementary Information, section 3) and the concentration versus depth composite profiles for both  $^{26}\text{Al}$  and  $^{10}\text{Be}$  reveal that the shallowest samples may have been exposed during a period of time (~15,000 years ago) that is consistent with deglaciation in the area (Supplemental Information). While we interpret the

individual simple burial age of all samples as a maximum limiting age of deposition of the Kap København Formation Member B, we recommend using the three most deeply shielded samples in a single depth profile to minimize the effect of post-depositional production. We then calculate a convolved probability distribution age for these three samples (KK06A, B and C). However, this calculation depends on the  $^{26}\text{Al}:\text{Be}$  production ratio we use (that is, between 6.75 and 7.42) and on whether we adjust for erosion in the catchment. So, we repeat the convolved probability distribution function age for the lowest and highest production ratio and zero to maximum possible erosion rate, to obtain the minimum and maximum limiting age range at 1 $\sigma$  confidence (Supplementary Information, section 3). Taking the midpoint between the negative and positive 3 $\sigma$  confidence limits, we obtain a maximum burial age of  $2.70 \pm 0.46$  Myr. This age is also supported by the position of those three samples on the isochron plot, which suggests the true age may not be significantly different than this maximum limiting age.

#### Thermal age

The extent of thermal degradation of the Kap København DNA was compared to the DNA from the Krestovka Mammoth molar. Published kinetic parameters for DNA degradation<sup>60</sup> were used to calculate the relative rate difference over a given interval of the long-term temperature record and to quantify the offset from the reference temperature of  $10^{\circ}\text{C}$ , thus estimating the thermal age in years at  $10^{\circ}\text{C}$  for each sample (Supplementary Information, section 4). The mean annual air temperature (MAT) for the the Kap København sediment was taken from Funder et al. (2001)<sup>61</sup> and for the Krestovka Mammoth the MAT was calculated using temperature data from the Cerskij Weather Station (WMO no. 251230)  $68.80^{\circ}\text{N}$   $161.28^{\circ}\text{E}$ , 32 m from the IRI Data Library (<https://iri.columbia.edu/>) (Supplementary Table 4.4.1).

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We did not correct for seasonal fluctuation for the thermal age calculation of the Kap København sediments or from the Krestovka Mammoth. We do provide theoretical average fragment length for four different thermal scenarios for the DNA in the Kap København sediments (Supplementary Table 4.4.2). A correction in the thermal age calculation was applied for altitude using the environmental lapse rate ( $6.49^{\circ}\text{C km}^{-1}$ ). We scaled the long-term temperature model of Hansen et al. (2013)<sup>62</sup> to local estimates of current MATs by a scaling factor sufficient to account for the estimates of the local temperature decline at the last glacial maximum and then estimated the integrated rate using an Ea of  $127\text{ kJ mol}^{-1}$  (ref. <sup>60</sup>).

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#### Mineralogic composition

The minerals in each of the Kap København sediment samples were identified using X-ray diffraction and their proportions were quantified using Rietveld refinement. The samples were homogenized by grinding ~1 g of sediment with ethanol for 10 min in a McCrone Mill. The samples were dried at  $60^{\circ}\text{C}$  and added corundum (CR-1, Baikowski) as the internal standard to a final concentration of 20.0 wt%. Diffractograms were collected using a Bruker D8 Advance ( $\Theta-\Theta$  geometry) and the LynxEye detector (opening 2.71°), with  $\text{Cu } K_{\alpha1,2}$  radiation (1.54 Å; 40 kV, 40 mA) using a Ni-filter with thickness of 0.2 mm on the diffracted beam and a beam knife set at 3 mm. We scanned from  $5\text{--}90^{\circ}\text{ 2}\theta$  with a step size of 0.1° and a step time of 4 s while the sample was spun at 20 rpm. The opening of the divergence slit was 0.3° and of the antiscatter slit 3°. Primary and secondary Soller slits had an opening of 2.5° and the opening of the detector window was 2.71°. For the Rietveld analysis, we used the Profex interface for the BGMIN software<sup>62,63</sup>. The instrumental parameters and peak broadening were determined by the fundamental parameters ray-tracing procedure<sup>64</sup>. A detailed description of identification of clay minerals can be found in the supporting information.

#### Adsorption

We used pure or purified minerals for adsorption studies. The minerals used and treatments for purifying them are listed in Supplementary

**Table 4.2.6.** The purity of minerals was checked using X-ray diffraction with the same instrumental parameters and procedures as listed in the above section i.e., mineralogical composition. Notes on the origin, purification and impurities can be found in the supplementary information section 4. We used artificial seawater<sup>65</sup> and salmon sperm DNA (low molecular weight, lyophilized powder, Sigma Aldrich) as a model for eDNA adsorption. A known amount of mineral powder was mixed with seawater and sonicated in an ultrasonic bath for 15 min. The DNA stock was then added to the suspension to reach a final concentration between 20–800 µg ml<sup>-1</sup>. The suspensions were equilibrated on a rotary shaker for 4 h. The samples were then centrifuged and the DNA concentration in the supernatant determined with UV spectrometry (Biophotometer, Eppendorf), with both positive and negative controls. All measurements were done in triplicates, and we made five to eight DNA concentrations per mineral. We used Langmuir and Freundlich equations to fit the model to the experimental isotherm and to obtain adsorption capacity of a mineral at a given equilibrium concentration.

#### Pollen

The pollen samples were extracted using the modified Grischuk protocol adopted in the Geological Institute of the Russian Academy of Science which utilizes sodium pyrophosphate and hydrofluoric acid<sup>66</sup>. Slides prepared from 6 samples were scanned at 400× magnification with a Motic BA 400 compound microscope and photographed using a Moticam 2300 camera. Pollen percentages were calculated as a proportion of the total palynomorphs including the unidentified grains. Only 4 of the 6 samples yielded terrestrial pollen counts ≥50. In these, the total palynomorphs identified ranged from 225 to 71 (mean = 170.25; median = 192.5). Identifications were made using several published keys<sup>67,68</sup>. The pollen diagram was initially compiled using Tilia version 1.5.12<sup>69</sup> but replotted for this study using Psimpoll 4.10<sup>70</sup>.

#### DNA recovery

For recovery calculation, we saturated mineral surfaces with DNA. For this, we used the same protocol as for the determination of adsorption isotherms with an added step to remove DNA not adsorbed but only trapped in the interstitial pores of wet paste. This step was important because interstitial DNA would increase the amount of apparently adsorbed DNA and overestimate the recovery. To remove trapped DNA after adsorption, we redispersed the minerals in seawater. The process of redispersing the wet paste in seawater, ultracentrifugation and removal of supernatant lasted less than 2.5 min. After the second centrifugation, the wet pastes were kept frozen until extraction. We used the same extraction protocol as for the Kap København sediments. After the extraction, the DNA concentration was again determined using UV spectrometry.

#### Metagenomes

A total of 41 samples were extracted for DNA<sup>71</sup> and converted to 65 dual indexed Illumina sequencing libraries (including 13 negative extraction and library controls)<sup>30</sup>. 34 libraries were thereafter subjected to ddPCR using a QX200 AutoDG Droplet Digital PCR System (Bio-Rad) following manufacturer's protocol. Assays for ddPCR include a P7 index primer (5'-AGCAGAAGACGGCATAC-3') (900nM), gene-targeting primer (900 nM), and a gene-targeting probe (250nM). We screened for Viridiplantae psbD (primer: 5'-TCATAATTGGACGTTAACCC-3', probe: 5'-(FAM)ACTCCCCATCATATGAAA(BHQ1)-3') and Poaceae psbA (primer: 5'-CTCACAACTTCCCTCTAGAC-3', probe 5'-(HEX) AGCTGCTTGAAGTTC(BHQ1)-3'). Additionally, 34 of the 65 libraries were enriched using targeted capture enrichment, for mammalian mitochondrial DNA using the PaleoChip Arctic1.0 bait-set<sup>31</sup> and all libraries were hereafter sequenced on an Illumina HiSeq 4000 80 bp PE or a NovaSeq 6000 100 bp PE. We sequenced a total of 16,882,114,068 reads which, after low complexity filtering (Dust =1), quality trimming ( $q \geq 25$ ), duplicate removal and filtering for reads longer than 29 bp

(only paired read mates for NovaSeq data) resulted in 2,873,998,429 reads that were parsed for further downstream analysis. We next estimated kmer similarity between all samples using simka<sup>32</sup> (setting heuristic count for max number of reads (-max-reads 0) and a kmer size of 31 (-kmer-size 31)), and performed a principal component analysis (PCA) on the obtained distance matrix (see Supplementary Information, 'DNA'). We hereafter parsed all QC reads through HOLI<sup>33</sup> for taxonomic assignment. To increase resolution and sensitivity of our taxonomic assignment, we supplemented the RefSeq (92 excluding bacteria) and the nucleotide database (NCBI) with a recently published Arctic-boreal plant database (PhyloNorway) and Arctic animal database<sup>34</sup> as well as searched the NCBI SRA for 139 genomes of boreal animal taxa (March 2020) of which 16 partial/full genomes were found and added (Source Data 1, sheet 4) and used the GTDB microbial database version 95 as decoy. All alignments were hereafter merged using samtools and sorted using gz-sort (v. 1). Cytosine deamination frequencies were then estimated using the newly developed metaDMG, by first finding the lowest common ancestor across all possible alignments for each read and then calculating damage patterns for each taxonomic level (<https://metadmg-dev.github.io/metaDMG-core/index.html>) (Supplementary Information, section 6). In parallel, we computed the mean read length as well as number of reads per taxonomic node (Supplementary Information, section 6). Our analysis of the DNA damage across all taxonomic levels pointed to a minimum filter for all samples at all taxonomic levels with a D-max ≥ 25% and a likelihood ratio (λ-LR) ≥ 1.5. This ensured that only taxa showing ancient DNA characteristics were parsed for downstream profiling and analysis and resulted in no taxa within any controls being found (Supplementary Information, section 6).

#### Marine eukaryotic metagenome

We sought to identify marine eukaryotes by first taxonomically labelling all quality-controlled reads as Eukaryota, Archaea, Bacteria or Virus using Kraken 2<sup>72</sup> with the parameters '--confidence 0.5 --minimum-hit-groups 3' combined with an extra filtering step that only kept those reads with root-to-leaf score >0.25. For the initial Kraken 2 search, we used a coarse database created by the taxdb-integration workflow (<https://github.com/AMG-tk/taxdb-integration>) covering all domains of life and including a genomic database of marine planktonic eukaryotes<sup>73</sup> that contain 683 metagenome-assembled genomes (MAGs) and 30 single-cell genomes (SAGs) from *Tara Oceans*<sup>74</sup>, following the naming convention in Delmont et al.<sup>73</sup>, we will refer to them as SMAGs. Reads labelled as root, unclassified, archaea, bacteria and virus were refined through a second Kraken 2 labelling step using a high-resolution database containing archaea, bacteria and virus created by the taxdb-integration workflow. We used the same Kraken 2 parameters and filtering thresholds as the initial search. Both Kraken 2 databases were built with parameters optimized for the study read length (-kmer-len 25 --minimizer-len 23 --minimizer-spaces 4).

Reads labelled as eukaryota, root and unclassified were hereafter mapped with Bowtie2<sup>75</sup> against the SMAGs. We used MarkDuplicates from Picard (<https://github.com/broadinstitute/picard>) to remove duplicates and then we calculated the mapping statistics for each SMAG in the BAM files with the filterBAM program (<https://github.com/AMG-tk/bam-filter>). We furthermore estimated the postmortem damage of the filtered BAM files with the Bayesian methods in metaDMG and selected those SMAGs with a D-max ≥ 0.25 and a fit quality (λ-LR) higher than 1.5. The SMAGs with fewer than 500 reads mapped, a mean read average nucleotide identity (ANI) of less than 93% and a breadth of coverage ratio and coverage evenness of less than 0.75 were removed. We followed a data-driven approach to select the mean read ANI threshold, where we explored the variation of mapped reads as a function of the mean read ANI values from 90% to 100% and identified the elbow point in the curve (Supplementary Fig. 6.11.I). We used anvi'o<sup>76</sup> in manual mode to plot the mapping and damage results using the SMAGs phylogenomic tree inferred by Delmont et al.

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as reference. We used the oceanic signal of Delmont et al. as a proxy to the contemporary distribution of the SMAGs in each ocean and sea (Fig. 5 and Supplementary Information, section 6).

### Comparison of DNA, macrofossil and pollen

To allow comparison between records in DNA, macrofossil and pollen, the taxonomy was harmonized following the Pan Arctic Flora checklist<sup>4</sup> and NCBI. For example, since Bennike (1990)<sup>18</sup>, *Potamogeton* has been split into *Potamogeton* and *Stuckenia*, *Polygonum* has been split to *Polygonum* and *Bistorta*, and *Saxifraga* was split to *Saxifraga* and *Micranthes*, whereas others have been merged, such as *Melandrium* with *Silene*<sup>39</sup>. Plant families have changed names—for instance, Gramineae is now called Poaceae and Scrophulariaceae has been re-circumscribed to exclude Plantaginaceae and Orobanchaceae<sup>77</sup>. We then classified the taxa into the following: category 1 all identical genus recorded by DNA and macrofossils or pollen, category 2 genera recorded by DNA also found by macrofossils or pollen including genus contained within family level classifications, category 3 taxa only recorded by DNA, category 4 taxa only recorded by macrofossils or pollen (Source Data 1).

### Phylogenetic placement

We sought to phylogenetically place the set of ancient taxa with the most abundant number of reads assigned, and with a sufficient number of reference sequences to build a phylogeny. These taxa include reads mapped to the chloroplast genomes of the flora genera *Salix*, *Populus* and *Betula*, and to the mitochondrial genomes of the fauna families Elephantidae, Cricetidae, Leporidae, as well as the subfamilies Capreolinae and Anserinae. Although the evolution of the chloroplast genome is somewhat less stable than that of the plant mitochondrial genome, it has a faster rate of evolution, and is non-recombining, and hence is more likely to contain more informative sites for our analysis than the plant mitochondria<sup>78</sup>. Like the mitochondrial genome, the chloroplast genome also has a high copy number, so that we would expect a high number of sedimentary reads mapping to it.

For each of these taxa, we downloaded a representative set of either whole chloroplast or whole mitochondrial genome fasta sequences from NCBI Genbank<sup>79</sup>, including a single representative sequence from a recently diverged outgroup. For the *Betula* genus, we also included three chloroplast genomes from the PhyloNorway database<sup>34,80</sup>. We changed all ambiguous bases in the fasta files to N. We used MAFFT<sup>81</sup> to align each of these sets of reference sequences, and inspected multiple sequence alignments in NCBI MSAViewer to confirm quality<sup>82</sup>. We trimmed mitochondrial alignments with insufficient quality due to highly variable control regions for Leporidae, Cricetidae and Anserinae by removing the d-loop in MegaX<sup>83</sup>.

The BEAST suite<sup>48</sup> was used with default parameters to create ultrametric phylogenetic trees for each of the five sets of taxa from the multiple sequence alignments (MSAs) of reference sequences, which were converted from Nexus to Newick format in Figtree (<https://github.com/rambaut/figtree>). We then passed the multiple sequence alignments to the Python module AlignIO from BioPython<sup>84</sup> to create a reference consensus fasta sequence for each set of taxa. Furthermore, we used SNPSites<sup>85</sup> to create a vcf file from each of the MSAs. Since SNPSites outputs a slightly different format for missing data than needed for downstream analysis, we used a custom R script to modify the vcf format appropriately. We also filtered out non-biallelic SNPs.

From the damage filtered ngsLCA output, we extracted all readIDs uniquely classified to reference sequences within these respective taxa or assigned to any common ancestor inside the taxonomic group and converted these back to fastq files using seqtk (<https://github.com/lh3/seqtk>). We merged reads from all sites and layers to create a single read set for each respective taxon. Next, since these extracted reads were mapped against a reference database including multiple sequences from each taxon, the output files were not on the same coordinate system. To circumvent this issue and avoid mapping bias, we

re-mapped each read set to the consensus sequence generated above for that taxon using bwa<sup>86</sup> with ancient DNA parameters (bwa aln -n 0.001). We converted these reads to bam files, removed unmapped reads, and filtered for mapping quality > 25 using samtools<sup>87</sup>. This produced 103,042, 39,306, 91,272, 182 and 129 reads for *Salix*, *Populus*, *Betula*, Elephantidae and Capreolinae, respectively. Q24

We next used pathPhynder<sup>88</sup>, a phylogenetic placement algorithm that identifies informative markers on a phylogeny from a reference panel, evaluates SNPs in the ancient sample overlapping these markers, and traverses the tree to place the ancient sample according to its derived and ancestral SNPs on each branch. We used the transversions-only filter to avoid errors due to deamination, except for *Betula*, *Salix* and *Populus* in which we used no filter due to sufficiently high coverage. Last, we investigated the pathPhynder output in each taxon set to determine the phylogenetic placement of our ancient samples (see supplementary information for discussion on phylogenetic placement).

Based on the analysis described above we further investigated the phylogenetic placement within the genus *Mammut*, or mastodons. To avoid mapping reference biases in the downstream results, we first built a consensus sequence from all comparative mitochondrial genomes used in said analysis and mapped the reads identified in ngsLCA as Elephantidae to the consensus sequence. Consensus sequences were constructed by first aligning all sequences of interest using MAFFT<sup>81</sup> and taking a majority rule consensus base in Geneious v2020.0.5 (<https://www.geneious.com>). We performed three analyses for phylogenetic placement of our sequence: (1) Comparison against a single representative from each Elephantidae species including the sea cow (*Dugong dugon*) as outgroup, (2) Comparison against a single representative from each Elephantidae species, and (3) Comparison against all published mastodon mitochondrial genomes including the Asian elephant as outgroup.

For each of these analyses we first built a new reference tree using BEAST v1.10.4 (ref. <sup>46</sup>) and repeated the previously described pathPhynder steps, with the exception that the pathPhynder tree path analysis for the *Mammut* SNPs was based on transitions and transversions, not restricting to only transversions due to low coverage.

***Mammut americanum*.** We confirmed the phylogenetic placement of our sequence using a selection of Elephantidae mitochondrial reference sequences, GTR+G, strict clock, a birth-death substitution model, and ran the MCMC chain for 20,000,000 runs, sampling every 20,000 steps. Convergence was assessed using Tracer<sup>89</sup> v1.7.2 and an effective sample size (ESS) > 200. To determine the approximate age of our recovered mastodon mitogenome we performed a molecular dating analysis with BEAST<sup>46</sup> v1.10.4. We used two separate approaches when dating our mastodon mitogenome, as demonstrated in a recent publication<sup>90</sup>. First, we determined the age of our sequence by comparing against a dataset of radiocarbon-dated specimens ( $n = 13$ ) only. Secondly, we estimated the age of our sequence including both molecularly ( $n = 22$ ) and radiocarbon-dated ( $n = 13$ ) specimens using the molecular dates previously determined<sup>90</sup>. We utilized the same BEAST parameters as Karpinski et al.<sup>90</sup> and set the age of our sample with a gamma distribution (5% quantile:  $8.72 \times 10^4$ , Median:  $1.178 \times 10^6$ ; 95% quantile:  $5.093 \times 10^6$ ; initial value: 74,900; shape: 1; scale: 1,700,000). In short, we specified a substitution model of GTR+G4, a strict clock, constant population size, and ran the Markov Chain Monte Carlo chain for 50,000,000 runs, sampling every 50,000 steps. Convergence of the run was again determined using Tracer.

### Molecular dating methods

In this section, we describe molecular dating of the ancient birch (*Betula*) chloroplast genome using BEAST v1.10.4 (ref. <sup>46</sup>). In principle, the genera *Betula*, *Populus* and *Salix* had both sufficiently high chloroplast genome coverage (with mean depth  $24.16 \times$ ,  $57.06 \times$  and

27.04×, respectively, although this coverage is highly uneven across the chloroplast genome) and enough reference sequences to attempt molecular dating on these samples. Notably, this is one of the reasons we included a recently diverged outgroup with a divergence time estimate in each of these phylogenetic trees. However, our *Populus* sample clearly contained a mixture of different species, as seen from its inconsistent placement in the pathPhynder output. In particular, there were multiple supporting SNPs to both *Populus balsamifera* and *Populus trichocarpa*, and both supporting and conflicting SNPs on branches above. Furthermore, upon inspection, our *Salix* sample contained a surprisingly high number of private SNPs which is inconsistent with any ancient or even modern age, especially considering the number of SNPs assigned to the edges of the phylogenetic tree leading to other *Salix* sequences. We are unsure what causes this inconsistency but hypothesize that our *Salix* sample is also a mixed sample, containing multiple *Salix* species that diverged from the same placement branch on the phylogenetic tree at different time periods. This is supported by looking at all the reads that cover these private SNP sites, which generally appear to be from a mixed sample, with reads containing both alternate and reference alleles present at a high proportion in many cases. Alternatively, or potentially jointly in parallel, this could be a consequence of the high number of nuclear plastid DNA sequences (NUPTs) in *Salix*<sup>91</sup>. Because of this, we continued with only *Betula*.

First, we downloaded 27 complete reference *Betula* chloroplast genome sequences and a single *Alnus* chloroplast genome sequence to use as an outgroup from the NCBI Genbank repository, and supplemented this with three *Betula* chloroplast sequences from the PhyloNorway database generated in a recent study<sup>29</sup>, for a total of 31 reference sequences. Since chloroplast sequences are circular, downloaded sequences may not always be in the same orientation or at the same starting point as is necessary for alignment, so we used custom code (<https://github.com/miwipe/KapCopenhagen>) that uses an anchor string to rotate the reference sequences to the same orientation and start them all from the same point. We created a MSA of these transformed reference sequences with Mafft<sup>61</sup> and checked the quality of our alignment by eye in Seqtron<sup>92</sup> and NCBI MsaViewer. Next, we called a consensus sequence from this MSA using the BioAlign consensus function<sup>84</sup> in Python, which is a majority rule consensus caller. We will use this consensus sequence to map the ancient *Betula* reads to, both to avoid reference bias and to get the ancient *Betula* sample on the same coordinates as the reference MSA.

From the last common ancestor output in metaDMG<sup>93</sup>, we extracted read sets for all units, sites and levels that were uniquely classified to the taxonomic level of *Betula* or lower, with at a minimum sequence similarity of 90% or higher to any *Betula* sequence, using Seqtk<sup>94</sup>. We mapped these read sets against the consensus *Betula* chloroplast genome using BWA<sup>86</sup> with ancient DNA parameters (-o 2 -n 0.001 -t 20), then removed unmapped reads, quality filtered for read quality ≥25, and sorted the resulting bam files using samtools<sup>86</sup>. For the purpose of molecular dating, it is appropriate to consider these read sets as a single sample, and so we merged the resulting bam files into one sample using samtools. We used bcftools<sup>86</sup> to make an mpileup and call a vcf file, using options for haplidity and disabling the default calling algorithm, which can slightly biases the calls towards the reference sequence, in favour of a majority call on bases that passed the default base quality cut-off of 13. We included the default option using base alignment qualities<sup>95</sup>, which we found greatly reduced the read depths of some bases and removed spurious SNPs around indel regions. Lastly, we filtered the vcf file to include only single nucleotide variants, because we do not believe other variants such as insertions or deletions in an ancient environmental sample of this type to be of sufficiently high confidence to include in molecular dating.

We downloaded the gff3 annotation file for the longest *Betula* reference sequence, MG386368.1, from NCBI. Using custom R code<sup>96</sup>,

we parsed this file and the associated fasta to label individual sites as protein-coding regions (in which we labelled the base with its position in the codon according to the phase and strand noted in the gff3 file), RNA, or neither coding nor RNA. We extracted the coding regions and checked in Seqtron<sup>92</sup> and R that they translated to a protein alignment well (for example, no premature stop codons), both in the reference sequence and the associated positions in the ancient sequence. Though the modern reference sequence's coding regions translated to a high-quality protein alignment, translating the associated positions in the ancient sequence with no depth cut-off leads to premature stop codons and an overall poor quality protein alignment. On the other hand, when using a depth cut-off of 20 and replacing sites in the ancient sequence which did not meet this filter with N, we see a high-quality protein alignment (except for the N sites). We also interrogated any positions in the ancient sequence which differed from the consensus, and found that any suspicious regions (for example, with multiple SNPs clustered closely together spatially in the genome) were removed with a depth cut-off of 20. Because of this, we moved forward only with sites in both the ancient and modern samples which met a depth cut-off of at least 20 in the ancient sample, which consisted of about 30% of the total sites.

Next, we parsed this annotation through the multiple sequence alignment to create partitions for BEAST<sup>46</sup>. After checking how many polymorphic and total sites were in each, we decided to use four partitions: (1) sites belonging to protein-coding positions 1 and 2, (2) coding position 3, (3) RNA, or (4) non-coding and non-RNA. To ensure that these were high confidence sites, each partition also only included those positions which had at least depth 20 in the ancient sequence and had less than 3 total gaps in the multiple sequence alignment. This gave partitions which had 11,668, 5,828, 2,690 and 29,538 sites, respectively. We used these four partitions to run BEAST<sup>46</sup> v1.10.4, with unlinked substitution models for each partition and a strict clock, with a different relative rate for each partition. (There was insufficient information in these data to infer between-lineage rate variation from a single calibration). We assigned an age of 0 to all of the reference sequences, and used a normal distribution prior with mean 61.1 Myr and standard deviation 1.633 Myr for the root height<sup>47</sup>; standard deviation was obtained by conservatively converting the 95% HPD to z-scores. For the overall tree prior, we selected the coalescent model. The age of the ancient sequence was estimated following the overall procedures of Shapiro et al. (2011)<sup>97</sup>. To assess sensitivity to prior choice for this unknown date, we used two different priors, namely a gamma distribution metric towards a younger age (shape = 1, scale = 1.7); and a uniform prior on the range (0, 10 Myr). We also compared two different models of rate variation among sites and substitution types within each partition, namely a GTR+G with four rate categories, and base frequencies estimated from the data, and the much simpler Jukes Cantor model, which assumed no variation between substitution types nor sites within each partition. All other priors were set at their defaults. Neither rate model nor prior choice had a qualitative effect on results (Extended Data Fig. 10). We also ran the coding regions alone, since they translated correctly and are therefore highly reliable sites and found that they gave the same median and a much larger confidence interval, as expected when using fewer sites (Extended Data Fig. 10). We ran each Markov chain Monte Carlo for a total of 100 million iterations. After removing a burn-in of the first 10%, we verified convergence in Tracer<sup>89</sup> v1.7.2 (apparent stationarity of traces, and all parameters having an Effective Sample Size > 100). We also verified that the resulting MCC tree from TreeAnnotator<sup>46</sup> had placed the ancient sequence phylogenetically identically to pathPhynder<sup>88</sup> placement, which is shown in Extended Data Fig. 9. For our major results, we report the uniform ancient age prior, and the GTR+G<sub>4</sub> model applied to each of the four partitions. The associated XML is given in Source Data 3. The 95% HPD was (2.0172, 0.6786) for the age of the ancient *Betula* chloroplast sequence, with a median estimate of 1.323 Myr, as shown in Fig. 2.

## Article

### Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

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### Data availability

Raw sequence data is available through the ENA project accession PRJEB55522. Pollen counts are available through <https://github.com/miwipe/KapCopenhagen.git>. Source data are provided with this paper.

### Code availability

All code used is available at <https://github.com/miwipe/KapCopenhagen.git>.

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**Acknowledgements** We acknowledge support from the Carlsberg Foundation for logistics to carry-out two expeditions to Kap København in 2006 and 2012 (S. Funder, principal investigator for Carlsberg foundation grant to LongTerm and Kap København—the age). The fieldwork in 2016 was supported by a grant to N.K.L. from the Villum Foundation. E.W. and K.H.K. thank the Danish National Research Foundation (DNR) and the Lundbeck Foundation for providing long-term funds to develop the necessary DNA technology that eventually made it possible to retrieve environmental DNA from these ancient deposits in the Kap København Formation. M.W.P. acknowledges support from the Carlsberg Foundation (CF17-0275). K.K.S. and S.J. acknowledge support from VILLUM FONDEN (00025352). I.G.A. and E.C. have received funding from the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation programme (grant agreement no. 819192). B.D.S. acknowledges support from the Wellcome Trust programme in Mathematical Genomics and Medicine (WT220023). J.Å.K. was supported by the Carlsberg Foundation (CF20-0238). C.B. acknowledges ERC Advanced Award Diatomit (grant agreement no. 835067). J.C.G. was supported by Natural Science and Engineering Research Council of Canada-Discovery Grant 06785 and Canada Foundation for Innovation grant 21305. M.J.C. acknowledges support from the Danish National Research Foundation DNRF128. We thank G. Yang for cosmogenic isotope AMS target chemistry. S. Funder for introducing us to the Kap København Formation and generating much of the platform that enabled us to conduct our research; T. O. Delmont for providing data and guidance on the SMAGs analysis; Minik Rosing for providing talc minerals; T. B. Zunic for providing tremolite, orthoclase and chlorite; Z. Vardanyan for help with the DNA extractions and library build; and L. B. Levy and D. Skov for their help collecting samples in 2016. This work was prepared in part by LLNL under contract DE-AC52-07NA27344; LLNL-JRNL-830653. E.W. thanks St John’s College, Cambridge for providing him with a stimulating environment for scientific thoughts and discussion.

**Author contributions** K.H.K. and E.W. conceived the idea. K.H.K., M.W.P. and E.W. designed the study. K.H.K., A.M.Z.B., A.S.T., N.K.L. and E.W. provided samples, context and carried out fieldwork. M.W.P. undertook the DNA laboratory analysis and taxonomic profiling. M.W.P., B.D.S. and B.D.C. performed the phylogenetic placement with the supervision of M.S. and R.D. B.D.S., M.W.P. and B.D.C. performed the genetic dating with the supervision of R.D. and J.J.W. M.W.P., T.S.K. and C.S.M. conceived, designed and performed the DNA damage estimates. K.K.S. and S.J. conceived and designed the DNA–mineral aspects of the study, interpreted and wrote about the DNA–mineral data, and participated in the thermal age calculations. K.H.K., M.W.P., A.H.R., A.R., I.G.A. and E.W. undertook the floristic analysis and interpretations. K.K.K. performed cartography and GIS analysis. I.S. designed and carried out palaeomagnetic analysis and interpreted the results. J.C.G. prepared and analysed eight samples for cosmogenic <sup>26</sup>Al and <sup>10</sup>Be and interpreted their burial age. I.G.A., E.C. and Y.W. provided access to the PhyloNorway reference database, and gave input to the phylogenetic placement of the chloroplasts. A.S.T. counted pollen from the six additional samples. J.Å.K. supported sediment provenance evaluation. M.B. provided mineralogical data from North Greenland. C.D., M.R., M.E.J. and B.S. designed and carried out ddPCR based assays to detect and identify ancient plant DNA in samples. A.F.-G. contributed to the bioinformatic analysis of SMAGs and C.B. contributed to interpretation of marine metagenomic signals. M.J.C. contributed to the thermal age and DNA modelling. M.E.A. contributed to the DNA decay rate estimates. K.H.K., M.W.P., A.H.R. and E.W. interpreted the results and wrote the manuscript with contributions from K.K.S., S.J., A.R., B.D.S., B.D.C., I.G.A., J.C.G., I.S. and N.K.L., with inputs from the other authors.

**Competing interests** The authors declare no competing interests.

#### Additional information

**Supplementary information** The online version contains supplementary material available at <https://doi.org/10.1038/s41586-022-05453-y>.

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**Peer review information** Nature thanks the anonymous reviewer(s) for their contribution to the peer review of this work. Peer review reports are available.

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## B Explainable ML and Anaemia

The following pages contain the draft paper:

Christoffer C. Jørgensen, **Christian Michelsen**, Troels C. Petersen, Henrik Kehlet (2022), “*Gender-specific haemoglobin thresholds in relation to preoperative risk assessment in fast-track total hip and knee arthroplasty*”.

Based on the same data as used on Paper II, the paper uses the SHAP curves to understand the machine learning model. In particular, it compares the preoperative haemoglobin level in the patient with the risk-score for being resubmitted to the hospital within 30 days after the operation, stratified by sex and operation type (knee vs. hip replacement).

**Type of article: Science Letter**

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**Gender-specific haemoglobin thresholds in relation to preoperative risk assessment in fast-track total hip and knee arthroplasty.**

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**Short title:**

Preoperative anaemia and recovery in fast-track THA and TKA

Within the last decade there has been an increasing focus on anaemia, iron deficiency and transfusion strategies leading to the concept of “patient blood management” (PBM), aiming at reducing the need for blood transfusions by preoperative optimisation of haemoglobin (Hb) and iron-status and use of intra- and postoperative restrictive transfusion protocols [1].

When diagnosing preoperative anaemia most practitioners have adhered to the WHO guidelines which were developed in 1968 and use gender specific criteria of a Hb of  $< 130 \text{ g.l}^{-1}$  for men and  $< 12 \text{ g.l}^{-1}$  for women [2]. However, these thresholds are based on studies with less sophisticated laboratory and epidemiological techniques than presently available and are consequently under current revision [3].

Furthermore, it has been argued that the WHO definitions of anaemia may not apply to surgical patients, as the relative blood-loss is larger in women, potentially leading to increased risk of allogenic blood transfusions and morbidity when using a gender specific lower preoperative anaemia threshold [4-6].

In total hip (THA) and knee arthroplasty (TKA) it is internationally acknowledged that preoperative iron deficiency anaemia should be corrected by treatment with intravenous (i.v.) iron [7]. However, detailed knowledge of the Hb threshold to increase the risk of postoperative morbidity, indications for treatment and whether it differs in men and women is sparse. The aim of this secondary analysis was to investigate the influence of preoperative Hb level in a comprehensive machine-learning model aimed at identifying patients at “high-risk” of medical complications leading to either a length of hospital stay of  $> 4$  days or 30-days readmission after an established fast-track THA and TKA [8]. While the primary study focused on comparing potential benefits of an overall machine-learning model in preoperative risk-prediction [9], this secondary analysis focus specifically on the influence of preoperative Hb level per se and potential differences according to gender and age.

We used a well-defined cohort of elective fast-track THA and TKA patients and evaluated the effect of preoperative Hb-level on the machine-learning model by SHAP-analysis which evaluates the individual effect of the variables included in a machine-learning model [10]. Furthermore, we assessed the distribution of Hb-levels and increases in risk-profile according to gender and age.

From January 2017 to August 2017, we included 3913 patients with a median length of stay of 1 day. Mean preoperative Hb was 154.8 (SD:15.12) but lower in women (149.4 vs. 162 g.l<sup>-1</sup>: p<0.001) and there were 30.5% of women vs. 12.0% of men with a Hb of <130 g.l<sup>-1</sup> (p<0.001). SHAP-analysis demonstrated an immediate steep increase in the risk-score for medical complications with a preoperative Hb below 147.6 g.l<sup>-1</sup>, and irrespective of gender and age (figure 1). Finally, the median SHAP-value of Hb-level was 0.35 (IQR:) in the patients with a Hb-level below 147.6 g.l<sup>-1</sup>. These results remained consistent regardless of analysing THA and TKA separately (online Supporting Information Figure S1a+b).

Our analysis demonstrates that in a comprehensive machine-learning risk-model, the preoperative Hb threshold was the same in men and women for an increased risk of prolonged length of stay or readmissions due to medical issues after fast-track THA and TKA. The threshold value of 147.6 g.l<sup>-1</sup> is remarkably close to the 130 g.l<sup>-1</sup> suggested for men in the current WHO guideline. Thus, the results of our study support the current WHO threshold for anaemia in men, but importantly also for removing gender specific Hb criteria for preoperative anaemia in women, at least in elective THA and TKA. Furthermore, the influence of preoperative Hb level < 147.6 g.l<sup>-1</sup> was consistent regardless of age, supporting that the removal of gender specific criteria should apply to all patients. Finally, the effect of Hb level on the accumulated risk-score was clinically meaningful. Thus, figure 1, illustrates that preoperative Hb level contributed with SHAP-values of approximately 0.4 in patients with a Hb of

<147.6 g.l<sup>-1</sup>. This corresponds with about 50% increased odds of being a high-risk patient. In contrast, in those with Hb-levels >147.6 g.l<sup>-1</sup> the odds of being high-risk patients decreased with about 15%.

That gender specific Hb criteria may be inappropriate and need further consideration, has also been demonstrated in cardiac surgery, where women with a preoperative Hb of 120-129 g.l<sup>-1</sup> received more blood transfusions and had increased length of hospital stay compared to those with a Hb of >129 g.l<sup>-1</sup> [11]. That women with a preoperative Hb level of < 130 g.l<sup>-1</sup> may potentially benefit from iron-treatment prior to surgery was illustrated by a large study investigating preoperative Hb levels and iron deficiency in major elective surgery and finding similar incidence of iron deficiency in women with Hb < 130 g.l<sup>-1</sup> and < 120 g.l<sup>-1</sup> [12]. Our study has some limitations, including lack of information on perioperative blood-transfusions and potential use of preoperative i.v. iron. However, preoperative optimisation with i.v. iron was not standard in the participating departments, and even if some of the outcomes was due to transfusion-related morbidity it would not change the finding of similar SHAP-curves between men and women. Study strengths include well-established fast-track protocols, detailed data on comorbidity and patient outcomes, a complete follow-up, and use of a sophisticated machine-learning model.  
In conclusion, from a machine-learning model in fast-track THA and TKA, a Hb threshold of 146.7 g.l<sup>-1</sup> was found to increase risk of impaired recovery, regardless of gender or age, thus calling for re-evaluation of preoperative anaemia risk criteria in the elective surgical setting.

**Competing Interests**

The study was sponsored by a grant from the Novo Nordisk Foundation.

**Acknowledgements**

The authors would like to acknowledge the members of the Fast-track Hip and Knee Replacement Centre Collaborative group.

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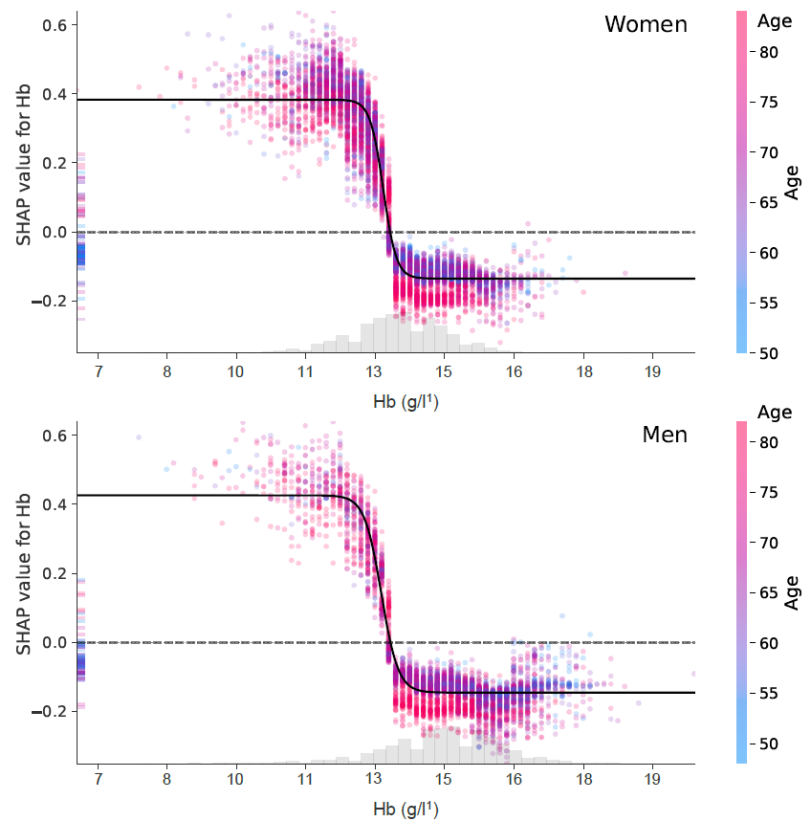
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**Figure legend:****Figure 1a+b**

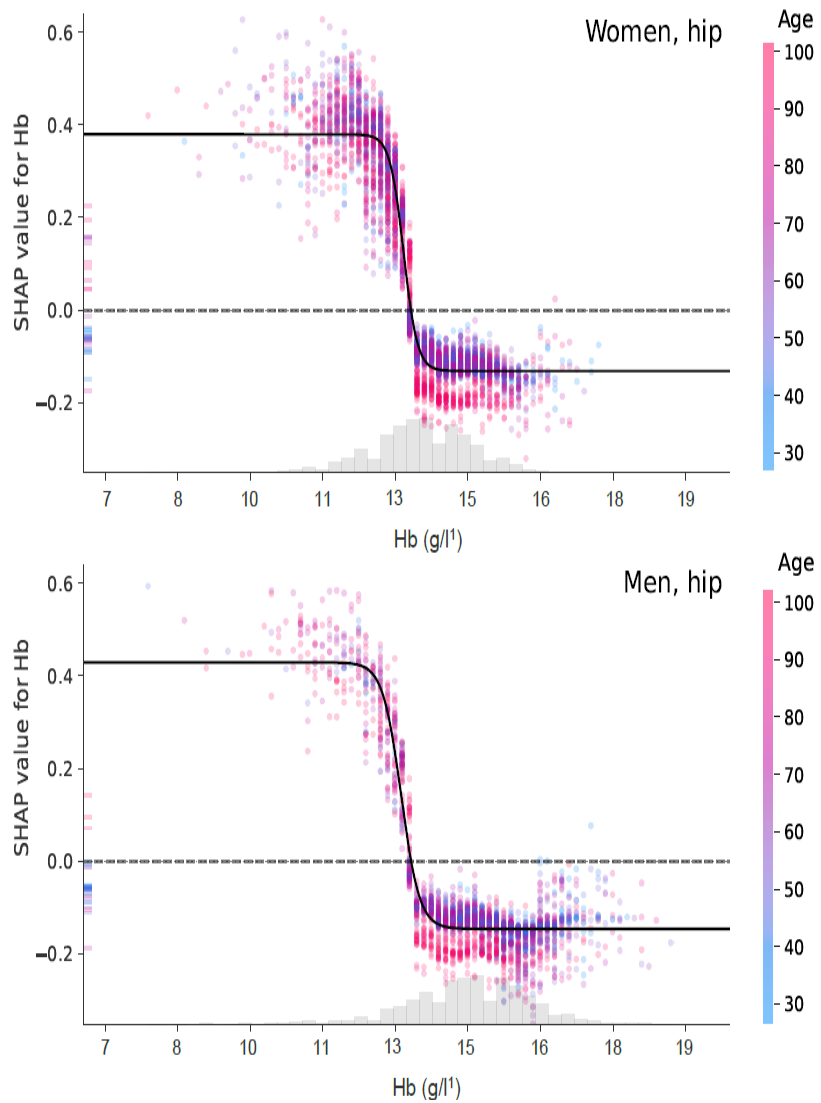
SHapley Additive exPlanations (SHAP) curves for preoperative haemoglobin level in relation to preoperative risk-stratification according to the machine-learning algorithm. Each dot indicates a patient with the colour indicating age (increasing from blue to red). Increasing SHAP values indicate increasing risk-score and decreasing values a decreased risk-score. The cut-off for going from a negative to a positive SHAP-value is indicated by the dotted line at a preoperative Hb level of 147.6 g.l<sup>1</sup>.

**Supplemental material****Figure 1a+b**

SHapley Additive exPlanations (SHAP) curves for preoperative haemoglobin level in relation to preoperative risk-stratification according to the machine-learning algorithm for total hip (1a) and total knee arthroplasty (1b), respectively. Each dot indicates a patient with the colour indicating age (increasing from blue to red). Increasing SHAP values indicate increasing risk-score and decreasing values a decreased risk-score. The cut-off for going from a negative to a positive SHAP-value is indicated by the dotted line at a preoperative Hb level of 147.6 g.l<sup>1</sup>.

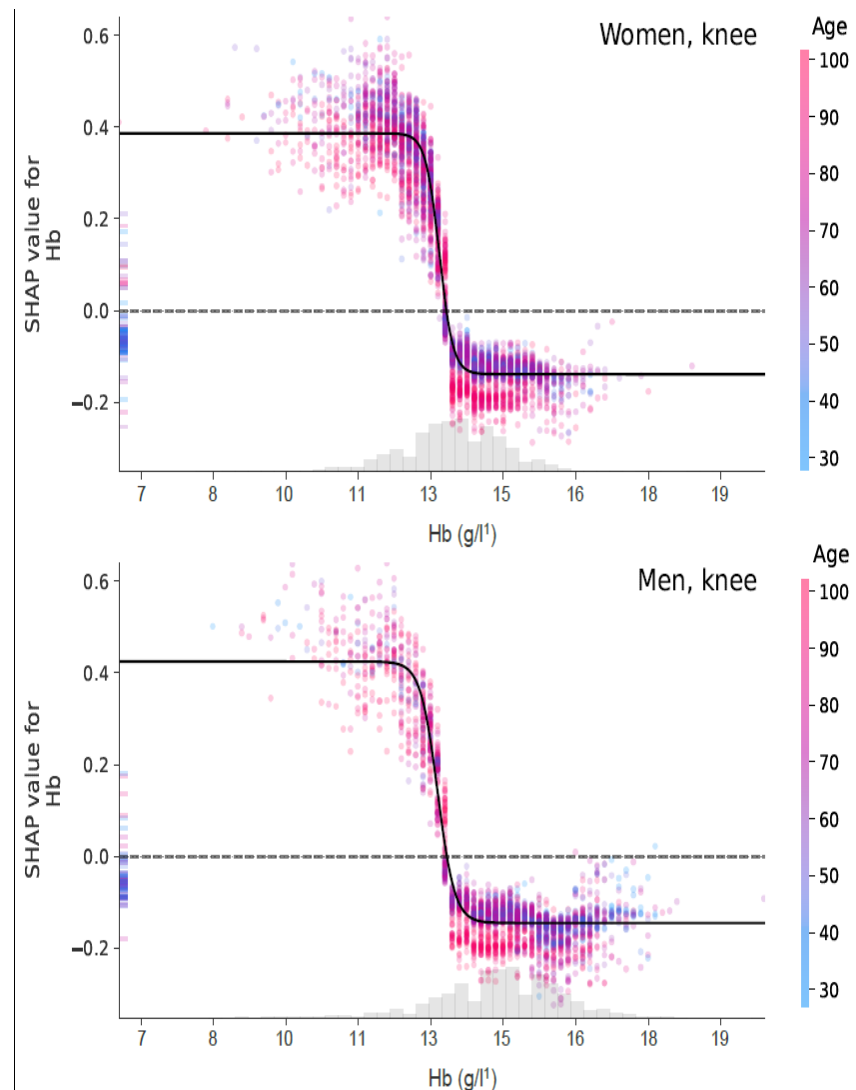
**Figure 1 a+b**

Supplemental figure 1a



10

Supplemental figure 1b





## C *SSI Ekspertrapport*

The following pages contain the report from Statens Serum Institut, the Danish CDC:

Ekspertgruppen for matematisk modellering, “*Ekspertrapport af den 10. december 2020 – Effekten af kontaktopsporing*” (Statens Serum Institut, 2020).

The report is from December 10, 2020 and is a summary on the effect of contact tracing related to COVID-19 in Denmark. The report is in Danish and is based on two agent based models, one from DTU and our model from NBI.

STATENS  
SERUM  
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# Ekspertrapport af den 10. december 2020

Effekten af kontaktopsporing



## Indhold

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## 1. Sammenfatning og konklusion

I indeværende rapport har modelgruppen for matematisk modellering af COVID-19 estimeret hvilke delelementer af kontaktopsporing, som er afgørende for at opnå størst mulig effekt af kontaktopsporing af nære kontakter til COVID-19 smittede personer.

Rapporten præsenterer resultater fra to forskellige agentbaserede modeller, som er udviklet af eksperter fra Danmarks Tekniske Universitet (DTU) og Københavns Universitet, Niels Bohr Institutet (NBI).

En agentbaseret model gør det muligt at modellere enkelte tiltag og deres effekt på smittespredningen af COVID-19. Forudsætningen for en præcis simulation er, at der er tilgængelige data, som kan informere modellen. Der er flere parametre, hvor der i nærværende arbejder er lavet antagelser på basis af de tilgængelige oplysninger. Det forventes, at nogle af disse kan belyses efterhånden som yderligere data frembringes. Hvor der ikke er specifikke eller komplette data, vil en agentbaseret model have unøjagtigheder eller risikere at være baseret på antagelser, som ikke nødvendigvis er retvisende. I modellerne anvendes der endvidere ens ventetidsfordelinger for alle agenter, selvom der i realiteten kan være lokale udsving i ventetider.

Sundhedsstyrelsen udkom d. 23. november 2020 med opdaterede retningslinjer for smitteopsporing af nære kontakter, herunder en udvidet definition af nære kontakter. Indevedende rapport er udviklet i henhold til de tidligere retningslinjer, og tager ikke højde for disse ændringer.

Der er i rapporten heller ikke taget højde for den stigende brug af private antigen test. Coronaopsporingen under STPS foretager også opsporing af nære kontakter, for primært tilfælde som er testet positiv for COVID-19 på sådanne antigen test.

### Konklusion

Modellerne peger på, at den største reduktion i kontakttallet kan nås ved effektiv opsporing for flest mulige primært tilfælde. Gevinsten i form af en reduktion i kontakttallet er således større, såfremt der sikres effektiv opsporing for samtlige primært tilfælde, relativt til reduktionen i kontakttallet, som kan opnås ved at nedbringe ventetiden til test og testsvar for primært tilfældet.

Ventetiden til test og testsvar for et primært tilfælde med COVID-19, har stor betydning for den reduktion af kontakttallet, som kan opnås gennem kontaktopsporing. De to uafhængigt udviklede modeller fra hhv. DTU og NBI finder begge, at for hver dag ventetiden til test og testsvar forsinkes for primære tilfælde, stiger kontakttallet med 4%. DTU-modellen finder endvidere, at ventetiden til et primært tilfælde booker en test og samtidig går i isolations har stor betydning for reduktionen i kontakttallet.

Modellerne viser endvidere, at med de anvendte ventetidsfordelinger, vil størstedelen af de nære kontakter som opspores, bliver testet så sent, at det er en mindre del af smitten, som forhindres. Det er derfor vigtigt at opspore nære kontakter hurtigst muligt efter eksponering, så de kan isoleres og blive testet på dag 4 og 6. Dette vil igen afhænge af den samlede ventetid til test og testsvar for primært tilfældet, som er forudsætningen for at opsporingen af nære kontakter kan initieres.

Den agentbaserede model fra NBI finder, at der er yderligere gevinst at hente ved at opspore nære kontakter i de netværk en person indgår i uden for husstand, job og skole. Det skyldes, at relativt få kontakter uden for husstand, job og skole opspores, og at disse kontakter ofte starter nye smittekedder i ikke ellers relaterede netværk. En bredere smitteopsporing har den fordel, at den potentielt finder de nye smittede, som ikke udviser symptomer.



## 2. Formål og baggrund

### 2.1 Formål og baggrund for modelgruppen

Statens Serum Institut indgår i det operationelle beredskab for smitsomme sygdomme og yder rådgivning og bistand til regeringen i forbindelse med den aktuelle pandemi. Som en del af denne opgave har Statens Serum Institut nedsat og leder en ekspertgruppe, der har til formål at udvikle matematiske modeller til at belyse udviklingen i COVID-19 i Danmark. Medlemmerne af ekspertgruppen fremgår af bilag 5.

Ekspertgruppens modellering var i foråret 2020 baseret på en populationsmodel, der har fokus på den gennemsnitlige adfærd i befolkningen. Populationsmodellen er bedst egnet, når udviklingen beskrives godt ved gennemsnittet. Derimod er populationsmodellen ikke det bedste værktøj til at beskrive de stokastiske hændelser i lokale udbrud, som aktuelt driver smittespredningen af COVID-19 i Danmark.

Siden sommeren 2020 har modelgruppen derfor udviklet to agentbaserede modeller, som er platformen for de analyser, modelgruppen forventes at levere i den kommende periode. De agentbaserede modeller kan, modsat en populationsmodel, estimere effekten ved enkelte tiltag, såsom effekten ved at nedbringe forsamlingsforbuddet, eller effekten af kontaktopsporing.

### 2.2 Formål med rapporten

Opsporingen af nære kontakter, foretaget af Styrelsen for Patientsikkerhed (STPS), er løbende udbygget i Danmark siden foråret 2020. Opgaven er vokset betydeligt i takt med, at det daglige antal nye COVID-19 tilfælde stiger, som følge af både en opblussen af epidemien, men også af, at testkapaciteten i Danmark er væsentligt udbygget hen over sommeren. Der testes aktuelt omkring 70.000 personer dagligt.

Formålet med denne rapport er at belyse, hvilke faktorer der er afgørende for at sikre en effektiv kontaktopsporing. Dette blyses ved at estimere effekten af centrale elementer i kontaktopsporringen, såsom ventetid til test og testresultat hos primærtildfældet, samt ventetid til at nære kontakter bliver opsporet og testet.



## 3. Opsporing og håndtering af nære kontakter i Danmark

### 3.1 Forudsætninger for en effektiv kontaktopsporing

Den vigtigste forudsætning for, at kontaktopsporing er et effektivt redskab til at nedbringe smitten med COVID-19 er, at der til hver en tid identificeres flest mulige smittede personer, som der derved kan udføres smitteopsporing for. Jo lavere mørketallet er, desto flere vil kunne smitteopspores. Det er derfor afgørende, at der sikres nem og hurtig adgang til test, først og fremmest for personer med COVID-19 lignende symptomer, men også for øvrige personer, der kunne have misstanke om at være smittet med COVID-19. Den Nationale Prævalensundersøgelse i Danmark har vist, at op mod 40-50% af dem, som havde antistoffer mod SARS-CoV-2 i blodet, ingen erindring havde om at have haft COVID-19 lignende sygdom<sup>1</sup>. Ved at udbyde adgang til test for flest mulige personer, vil man også finde flere asymptomatiske smittebærere.

### 3.2 Definition af en nær kontakt

*Sundhedsstyrelsen udkom d. 23. november 2020 med opdaterede retningslinjer for smitteopsporing af nære kontakter. Indenværende rapport er udviklet i henhold til de tidligere retningslinjer.*

*Der er således ikke taget højde for den udvidede definition af nære kontakter, eller indførslen af screeningsprøver for personer, som ikke umiddelbart opfylder kriteriet for nære kontakter, men som har været eksponeret i et omfang hvor der tilrådes en screeningstest.*

Kontaktopsporingen af nære kontakter baserer sig på, at personer der testes positiv for COVID-19 isolerer sig, og dernæst at nære kontakter til den smittede opspores, isoleres og testes, for der ved at afbryde smittekæder hurtigst muligt.

Definitionen af en nær kontakt er beskrevet i Sundhedsstyrelsens rapport om smitteopsporing af nære kontakter<sup>2</sup>.

En nær kontakt er defineret som en af følgende personer:

- En person der bor sammen med en, der har fået påvist COVID-19
- En person der har haft direkte fysisk kontakt (fx kram) med en, der har fået påvist COVID-19
- En person med ubeskyttet og direkte kontakt til smittefarlige sekreter fra en person der har fået påvist COVID-19
- En person der har haft tæt ”ansigt-til-ansigt” kontakt inden for en 1 meter i mere end 15 minutter (fx i samtale med personen) med en, der har fået påvist COVID-19
- Sundhedspersonale og andre som har deltaget i plejen af en patient med COVID-19, og som ikke har anvendt værnemidler på de forskrevne måder

<sup>1</sup> <https://www.ssi.dk/-/media/arkiv/dk/aktuelt/nyheder/2020/notat---covid-19-prvalensundersgelsen.pdf?la=da>

<sup>2</sup> <https://www.sst.dk/da/Udgivelser/2020/COVID-19-Smitteopsporing-af-naere-kontakter>



### 3.3 Periode for smitteopsporing

Der foretages smitteopsporing for perioden, hvor primærtilfældet vurderes at være smitsom. Smitteperioden er således afgrænset til 48 timer før symptomdebut til 48 timer efter symptomophør. For primære tilfælde der ikke har symptomer på COVID-19, er den smitsomme periode afgrænset til 48 timer før positiv test til 7 dage efter.

### 3.4 Opsporing af nære kontakter

Nære kontakter til en person der er smittet med COVID-19 kan opspores på følgende måder:

- De bliver kontaktet af STPS's Coronaopsporingen
- De bliver kontaktet ifm. kendte udbrud, eksempelvis på skoler
- De bliver kontaktet direkte af primærtilfældet
- De bliver notificeret om, at de har været tæt på en smittet person via appen Smitte|Stop

#### *Nære kontakter opsporet af Coronaopsporingen*

Coronaopsporingen under STPS kontakter smittede mhp. at hjælpe med at identificere og opspore nære kontakter til den smittede. Smittede kan også vælge selv at iværksætte opsporing af nære kontakter, og henvise dem til Coronaopsporingen, hvor de nære kontakter vil modtage rådgivning om, hvornår de bør testes, samt får adgang til at booke test på de pågældende dage.

Aktivitetsrapporter fra STPS viser, at der i hele opsporingsperioden i gennemsnit opspores ca. 5 nære kontakter for hvert primærtfalde, der foretages kontaktsporing for. Dette er et samlet gennemsnit for opsporede nære kontakter som STPS opsporer, og som primærtilfældet selv opsporer.

Til sammenligning er det estimeret i HOPE-projektet, at danskere henover sommeren i gennemsnit havde ca. 11 kontakter dagligt. Dette antal er nu faldet til ca. 7 kontakter dagligt, som opfylder kriterierne for en nær kontakt, se bilag 4.

Det skal dog pointeres, at Coronaopsporingen ikke er involveret i opsporing af nære kontakter i relation til udbrud i dagtilbud, skoler, plejehjem og hospitaler. Der vil der være opspored kontakter fra sådanne udbrud, som kontakter Coronaopsporingen for at få rådgivning om hvilke dage de bør testes, samt for at få rekvizitioner til booking af test på de pågældende dage.

Nære kontakter anbefales at blive testet på dag 4 og dag 6 efter vurderet eksponering. Dette relaterer sig til latentstiden, som er perioden fra, at man bliver smittet, til at man er smitsom, og virus kan påvises. En person som er opsporet som nær kontakt til en smittet skal ifølge Sundhedsstyrelsens retningslinjer gå i selv-isolation, indtil der foreligger testsvar. Såfremt der foreligger et negativt testresultat på dag 4, kan den nære kontakt bryde isolationen, men skal fortsat testes på dag 6. Hvis testresultatet på dag 4 derimod er positivt, skal den nære kontakt ikke testes igen på dag 6, men forblive i isolation indtil 48 timer efter symptomophør.

#### *Nære kontakter der ikke opspores af Coronaopsporingen*

Der vil være nære kontakter, der ikke opspores og rådgives via Coronaopsporingen. Dette kunne fx være nære kontakter, der bliver opsporet af primærtilfældet selv, og som vælger at booke test på coronaprover.dk uden først at have rådført sig med Coronaopsporingen. Det kunne også være personer, som er opsporet via appen Smitte|Stop, eller personer der mener, at de på anden vis



kan være nære kontakter til en smittet – uden nødvendigvis at opfylde kriteriet for at være en nær kontakt.

I oktober måned blev der i alt testet 1.091.966 personer. Heraf havde 62% (n = 675.623) bestilt tid på coronaprover.dk. Blandt disse svarede 58% (n = 391.146) på spørgeskemaet på coronaprover.dk, hvoraf 25% (n = 99.389) anførte, at de blev testet fordi, de var nær kontakt til en smittet (herunder personer som er adviseret af Smitte|Stop app). Kun 13% (n = 12.706) af dem som svarede, at de blev testet fordi de var nær kontakt til en smittet, var testet på én af de rekvisitionskoder, som der anvendes i Coronaopsporingen (Tabel 1). Samlet set blev 45.616 personer testet på én af de rekvisitionskoder som anvendes i Coronaopsporingen i oktober måned, hvor test-positivprocenten var ca. 4%. Til sammenligning var positivprocenten for de personer, der svarede, at de var nær kontakt til en smittet på Coronaprover.dk omkring 2,5 %. Dette indikerer at Coronaopsporingen har større succes med at opspore de korrekte nære kontakter, sammenlignet med hvis befolkningen selv booker test som nær kontakt, uden forudgående rådgivning fra Coronaopsporingen.

*Tabel 1. Oversigt over antal testede i oktober måned 2020.*

	<b>Oktober</b>			
	<i>Testpositive (1. test)</i>	N	n	%
Testede personer	1.091.966	14.723	1.35	
Total antal tests rekvisiteret via Coronaopsporingen	45.616	1.941	4,26	
Bestilt på coronaprøver.dk	675.623	10.335	1,53	
Svaret på spørgeskema	391.146	5.387	1,38	
Ja, nær kontakt til smittet (herunder adviseret på Smitte Stop app)	99.389	2.544	2,56	
Rekvireret test via Coronaopsporingen	12.706	524	4,12	



## 4. Agentbaserede modeller

### 4.1 Om agentbaserede modeller

I indeværende rapport er resultaterne for effekten af kontaktopsporing frembragt fra to forskellige agentbaserede modeller, som er udviklet på henholdsvis Danmarks Tekniske Universitet (DTU) og Niels Bohr Institutet, Københavns Universitet (NBI).

En agentbaseret model simulerer et antal agenter (individer i en population) og deres interaktioner med andre agenter, svarende til de interaktioner som en befolkning normalt vis har. Hver agent er således en person, som er knyttet til en lokation i Danmark, svarende til deres bopæl., Agenterne indgår i flere forskellige netværk, f.eks. husstand, job og skole hvor de har kontakt til andre personer. Desuden har de andre kontakter til tilfældige personer i samfundet i den tid, hvor personen ikke er hjemme, på job eller i skole.

Hvis en agent bliver smittet med SARS-CoV-2, er forløbet for den enkelte agent beskrevet således, at agenten først er eksponeret (E) og derefter infektøs (I), hvorefter agenten ikke længere er smitsom og betragtes som rask (R). De gennemsnitlige tider i hvert sygdomsstadiu kan findes i bilag 1 og 2.

Hver kontakt som en agent eksponeres for tildeles en sandsynlighed for at blive smittet af en anden agent, hvis denne er smitsom. Sandsynligheden er sat til et niveau, som afspejler den nuværende situation med et kontakttal omkring 1.

Ud fra de ovenstående generelle antagelser simuleres en epidemi. For en mere detaljeret beskrivelse af de agentbaserede modeller, herunder de inkluderede parametre, henvises til bilag 1 (NBI) og 2 (DTU).

### 4.2 Forbehold

Mens en agentbaseret model kan medtage mere detaljerede dynamikker i en epidemi, så kræver en præcis simulation input fra data, som ofte ikke er tilgængelige eller forefindes, fx hvem en person mødes med i løbet af en dag. Derfor kan en sådan model have unøjagtigheder eller bygge på antagelser, som ikke er retvisende. Det er ikke muligt at kvantificere den nojagtige størrelse eller effekt af disse potentielle fejlkilder.



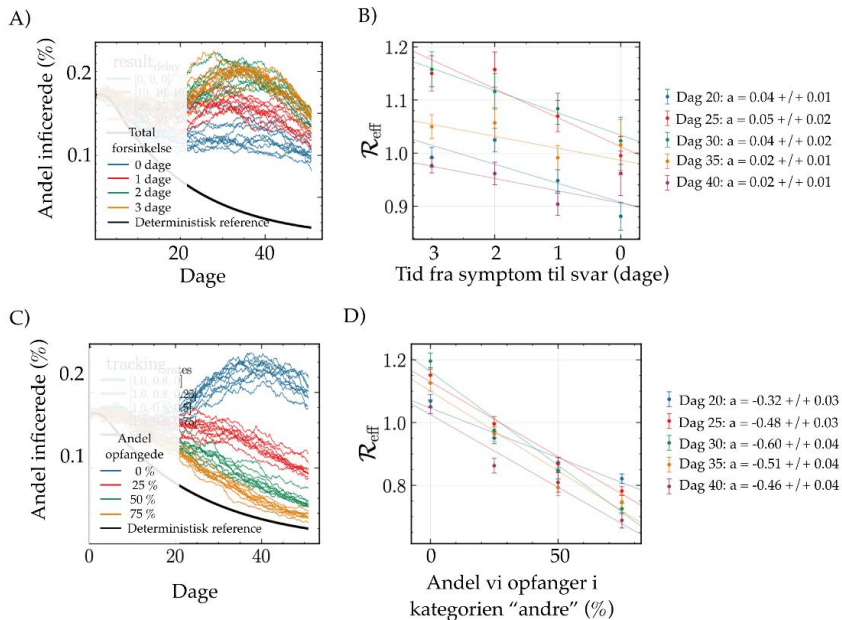
## 5. Resultater

### 5.1 Resultater fra den agentbaserede model udviklet af Niels Bohr Institutet, Københavns Universitet.

Modelkørslerne viser, at når 80% af de sekundære tilfælde i netværkenes husstande, arbejde og skole opspores, vurderes det, at ville nedsætte kontakttallet med omkring 30% sammenlignet med et hypotetisk scenarie uden opsporing af nære kontakter. Dette fremgår af figur 1. Hvis det af logistiske eller kapacitetsmæssige årsager ikke lykkedes at kontakte alle nye COVID-19 tilfælde, vil det betyde en forøgelse af kontakttallet i proportion til dette tal. Dvs. hvis opsporingen ikke kommer i kontakt med 20% af nye COVID-19 tilfælde, vil man potentielt miste 6 procentpoint ( $0.2 \times 0.3 = 0.06$ ) af reduktionen i kontakttallet, som ellers kunne opnås ved kontaktopsporing.

Ventetiden fra at et primært tilfælde ønsker en COVID-19 test (fx hvis man har symptomer), til at vedkommende har modtaget resultatet fra en test har indflydelse på effekten af både selvisolation og kontaktopsporing. Ved en række simulationer med forskellige antagelser finder modellen, at for hver dag man forkorter tiden mellem bestilling af test og testresultat mindskes kontakttallet med omkring 4%. Effekten er lidt større ved højere kontakttal end 1.

Effekten af kontaktopsporing kan øges ved at opspore flere i netværket af øvrige kontakter (ud over husstand og job og skole). Den agentbaserede model viser, at hvis man opsporer 25% af øvrige kontakter, vil kontakttallet falde med omkring 10%. En mere komplet kontaktopsporing (evt. yderligere hjulpet af apps på mobiltelefoner) vil således nedsætte kontakttallet væsentligt. Tilsvarende resultater er fundet i lignende modeller (Plank et al. (september 2020) og Kretzschmar et al. (august 2020)).



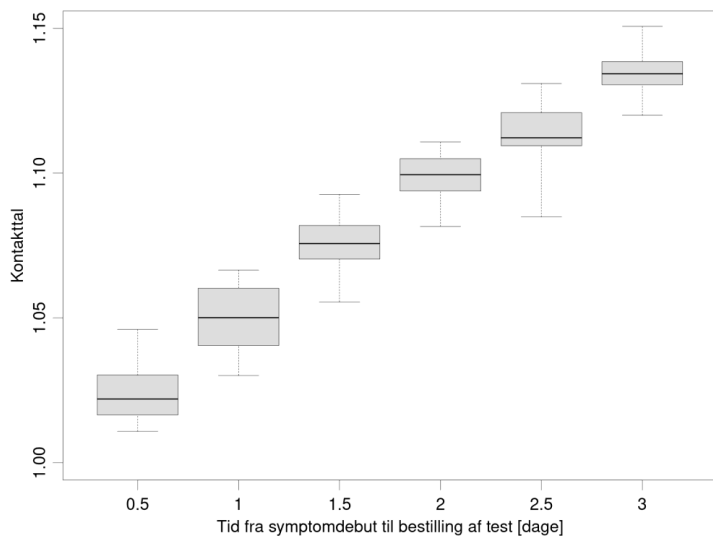


*Figur 1: A) Simuleret model, hvor hver kørsel (markeret med samme farve) gentages 10 gange for forskellige værdier af tiden fra symptom til svar. B) Værdien af kontakttallet estimeret på forskellige tidspunkter i simulationen vist i A). Den lineære sammenhæng giver en værdi for hvor mange procent kontakttallet sænkes for hver dag, man gør opsporingen hurtigere. C) Samme som A, men her for forskellige værdier af hvor mange man opsporer blandt øvrige kontakter D) Samme som B) men som funktion af hvor mange man opsporer blandt øvrige kontakter.*

### 5.2 Resultater fra den agentbaserede model udviklet af DTU Compute, Danmarks Tekniske Universitet

Denne agentbaserede model er baseret på tilhørsforhold til grupper (hjem, arbejdsplads, m.fl.). Modellen indeholder en række ventetider fra et primærtifælde får symptomer til sekundære tilfælde er opsporet. Modellen er nærmere beskrevet i bilag 2. Modellen er kørt med en række forskellige kombinationer af parametre. For hver kombination er der lavet 40 gentagelser for at illustrere variabiliteten. For hver gentagelse simuleres 30 dage som en transient, hvorefter kontakttallet estimeres baseret på de efterfølgende 30 dage.

De to parametre, som betyder mest for effekten af kontaktsporingen, er den gennemsnitlige ventetid fra en smittet får (milde) symptomer til at denne går i isolation og samtidig bestiller en test, samt andelen af kontakter som personen reducerer i perioden fra der bestilles en test til der foreligger et testsvar – det antages, at nære kontakter som opspores opretholder samme grad af isolation som andre, der venter på testsvar, hvilket vil sige, at nære kontakter går i isolation fra de bliver notificeret og indtil de får svar på deres første test.

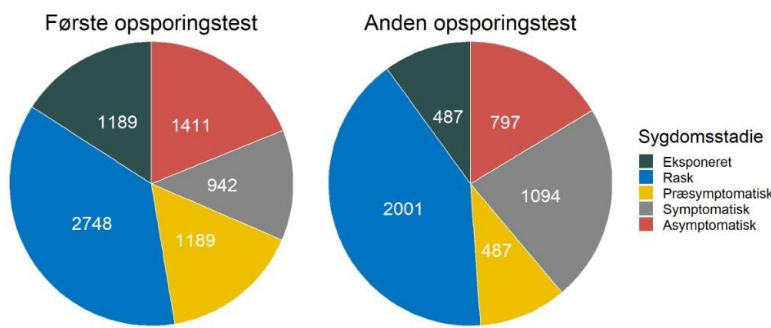


*Figur 2: Kontakttallets afhængighed af den gennemsnitlige tid fra at primærtifældet har symptomdebut til der bestilles en test og personen går i en grad af isolation. For hver parameterværdi er der foretaget 40 simulationer, og boxplottet viser median, de indre quartiler samt minimum og maksimum af disse.*



På figur 2 ses en klar effekt af tiden fra symptomdebut til isolation og samtidig bestilling af test. For hver dag den gennemsnitlige person går hurtigere i (delvis) isolation estimeres det, at kontakttallet reduceres med 0,04 (når referencen er et kontakttal omkring 1).

Modellen viser også, at omkring 25% af alle test positive, er fundet gennem kontaktopsporing. Det er her antaget, at der udføres kontaktopsporing for alle tilfælde (Se detaljer i bilag 2), samt at test af nære kontakter bestilles på de foreskrevne tidspunkter. Endvidere viser modellen, at over halvdelen af alle smittede aldrig bliver testet positiv (både falsk negative test og asymptotiske tilfælde). Disse starter derfor nye smittekæder uden forudgående opsporing. Dette kan være årsagen til, at det kun er 25% som findes gennem kontaktopsporing.



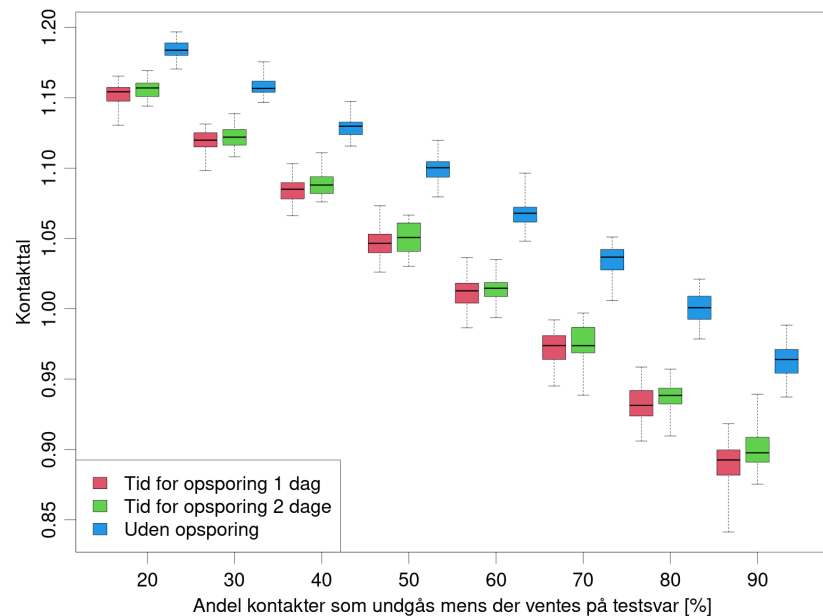
*Figur 3. Antal opsporede og smittede i hvert sygdomsstadije, når de får foretaget hhv. første og anden test i opsporingsprocessen. Der er flere, som ikke kommer tilanden test, bl.a. fordi de tester positiv i første test eller efter negativt testsvar vælger ikke at få taget den opfølgende test. Derudover vil der være en andel, hvor kontaktopsporingen er initieret sent, således at det kun er foreskrevet at teste personen én gang.*

Figur 3 beskriver de forskellige sygdomsstadier for smittede personer, som er opsporet som nære kontakter. Det ses, at en betydelig andel af de opsporedes personer, med de i modellen anvendte ventetidsfordelinger, på tidspunktet for opsporingen allerede har overstået deres infektionsperiode, når de testes første gang – en del af disse vil være smittet tidligere og ikke i forbindelse med den nærværende kontaktopsporing. I praksis vil nogle af disse teste positiv, da qPCR kan detektere virus 17 dage efter symptomdebut (Cevik et al., 2020). Desuden ses det, at personer i det præsymptomatiske stadije - hvor ca. halvdelen af smitten sker - kun udgør en lille andel af de opsporedes smittede personer ved både første og anden test. Ved begge test er det således under halvdelen af dem, som er smittede, som reelt er infektion. Kontaktopsporningen vil derfor kunne optimeres yderligere, hvis man identificerer flere nære kontakter i den præsymptomatiske fase. Dette kan ske ved at nedbringe ventetiden fra symptomdebut til testsvar for primærtildældet.

Personer, som tidligere er testet positiv er ikke medtaget her og bidrager derfor ikke til antallet af raske. Endvidere vil personer som modtager et positivt testresultat på deres første opsporingstest ikke få foretaget anden opsporingstest. Ovenstående diagrammer er produceret på baggrund af referenceparametrene som beskrevet i bilag 2.



Graden hvorved en smittet person isolerer sig, dvs. hvor stor en andel af ens kontakter man reducerer i perioden fra bestilling af test til testsvar, har stor betydning for kontakttallet. Referenceværdien antages at være 50% reduktion i antallet af kontakter i denne periode. Som det fremgår af figur 4 så opnås der i modellen en reduktion i kontakttallet på knap 0,04 for hver 10 procent-point graden af isolation øges, hvis der udføres kontaktopsporing (rød og grøn). Mens reduktionen er på 0,03 når der ikke udføres kontaktopsporing (blå). Således har andelen af kontakter, der reduceres hos primærtifældet og opsporedé nære kontakter i ventetiden fra bestilling af test til testsvar, større betydning for en reduktion i kontakttallet, end en reduktion i ventetiden til opsporing af nære kontakter.



Figur 4. Kontakttallets afhængighed af andelen af kontakter et primærtifælde og opsporedé nære kontakter reducerer, i ventetiden fra der bestilles en test til at testsvar foreligger, samt betydningen af ventetiden til at en nære kontakt opspores og går i tilsvarende isolation. For hver parameterværdi er der foretaget 40 simulationer og boxplottet viser median, de indre kvartiler samt minimum og maksimum af disse.



## 6. Referencer

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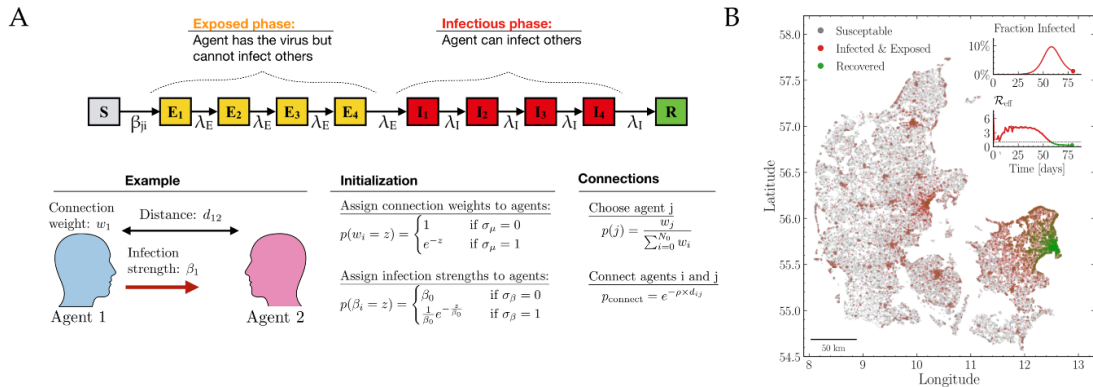
Plank, M., James, A., Lustig, A., Steyn, N., Binny, R. & Hendy, S. (2020). Potential reduction in transmission of COVID-19 by digital contact tracing systems. *MedRxiv*. Lokaliseret: <https://doi.org/10.1101/2020.08.27.20068346>



## Bilag 1. Beskrivelse af den agentbaserede model fra Niels Bohr Institutet

Bidrag og udvikling: Christian Michelsen, Emil Martiny, Tariq Halasa, Mogens H. Jensen, Troels C. Petersen og Mathias L. Heltberg

Den agentbaserede model fra NBI baseres på agenter, dvs. individer hvis karakteristika er tildelt ud fra statistiske fordelinger i befolkningen. Dette er f.eks. en aldersfordeling og en fordeling over pendlerafstande. Modellen starter med at fordele Danmarks bopæle ud i landet baseret på det danske hussalg over de sidste 15 år. Herefter placeres agenter i hver husstand baseret på deres alder og geografiske placering.



Figur 5: A) Skematisk oversigt over hvordan interaktionsnetværket i modellen ser ud. B) Eksempel på simulation af smittespredning i Danmark i modellen, gennem et simuleret tilfælde af flokimmunitet i København.

Et afgørende element i modellen er opbygningen af alle personers interaktionsnetværk. Dette genereres ved, at hver agent har et netværk, de interagerer med. Dette opdeles i tre dele: 1) kontakter i hjemmet, 2) kontakter på arbejdet, 3) kontakter i kategorien andre kontakter. Der er ikke nogen geografisk afhængighed af antallet af kontakter på arbejdet, men i den kategori der kaldes "andre", vil der generelt være flere kontakter for dem der bor i tæt befolkede områder i forhold til dem der bor på landet. Måden hvorpå netværket dannes er vist i Figur 5A.

Ud fra data fra HOPE-projektet har vi estimeret, hvor mange personer hver agent vil interagere med, og i denne model vil alle agenter have mellem 3 og 15 daglige kontakter.

Når modellen simuleres vil alle inficerede agenter gennemgå et forløb, hvor de er i en latent periode, hvor de ikke smitter, hvorefter de vil rykke over i en infektiøs periode, hvor de kan smitte agenter i deres netværk. Denne model simuleres ud fra det der kaldes Gillespie algoritmen, således at netværket opdateres instantant for alle smittebegivenheder. En samling af de væsentligste parametre er vist herunder (Tabel 2).



Tabel 2: Parametre i modellen.

Parameter	Værdi interval for middel-værdien	Reference
Antal kontakter per dag	3-15	HOPE projektet
Latent tid (dage)	3-5	Litteratur se referenceliste i bilag 5
Infektios tid (dage)	4-8	Litteratur se referenceliste i bilag 5
Andel af kontakter i "andre" (%)	30-80	HOPE projektet
Typisk afstand mellem kontakter (km)	5-20	Trafik data
Andel afstandsuafhængige kontakter (%)	3-5	Trafik data
Tid fra symptom til test (Dage)	0-2	Fordeling fra spørgeskemaundersøgelse i foråret 2020 (ikke offentliggjort)
Sandsynlighed for at få symptomer og blive testet (%)	20-60 %	Prævalensundersøgelsen
Sandsynlighed for at kontakte husstand (%)	100%	Antagelse
Sandsynlighed for at kontakte kollegaer (%)	40-80	Antagelse
Sandsynlighed for at kontakte andre (%)	0-75	Antagelse



## Bilag 2. Beskrivelse af den agentbaserede model fra DTU

*Bidrag og udvikling: Freja Terp Petersen, Jacob Bahnsen Schmidt, Kasper Telkamp Nielsen, Rebekka Quistgaard-Leth, Kaare Græsbøll og Lasse Engbo Christiansen*

Den agentbaserede model fra DTU baseres på en befolkningstabell, hvor hver række i tabellen svarer til en agent - eller et individ – og hver kolonne indeholder data, der beskriver den pågældende agent, herunder aldersgrupper med 5 års-intervaller, bopælskommune, netværks-ID og forskellige smitteparametere.

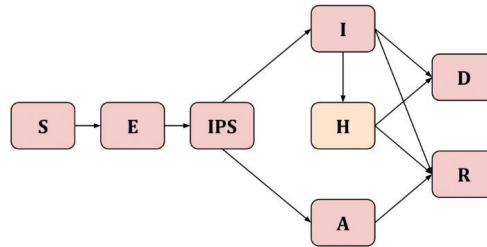
I sygdomsmodellen bæres smitten fremad ved, at agenter der deler netværks-ID, f.eks. husholdnings-ID, skole/job-ID eller omgangskreds-ID, kan smitte hinanden. Hver dag får alle agenter udregnet deres sandsynlighed for at blive smittet på baggrund af antal infektioner i deres forskellige netværk og på baggrund af deres individuelle antal nære kontakter, som de er blevet tildelt baseret på en fordeling fra totalt antal kontakter inden for 1m i HOPE projektet.

Der er 7 forskellige netværkstyper, som en agent kan være en del af:

- Husholdning (alle agenter har en husholdning)
- Daginistitution (børn mellem 0 og 4 år)
- Grundskole (børn mellem 5 og 14 år)
- Ungdomsuddannelse (unge mellem 15-24 år samt voksne på erhvervsuddannelser)
- Arbejdsplads med kontorinddelinger (voksne op til 65 år)
- Omgangskreds (alle agenter har en omgangskreds)
- Kommune (alle agenter har en kommune)

Agenterne er blevet tildelt netværk baseret på data fra Danmarks Statistik (husholdninger og arbejdspladser), Undervisningsministeriet (grundskoler og ungdomsuddannelser) samt Institution.dk (daginistitutioner).<sup>3</sup> Det antages i modellen, at den gennemsnitlige kontorstørrelse og den gennemsnitlige omgangskreds uden for skole og arbejde er på 8 personer.

<sup>3</sup> FAM122N: <https://www.statistikbanken.dk/FAM122N>  
 FAM133N: <https://statistikbanken.dk/FAM133N>  
 FAM55N: <https://statistikbanken.dk/FAM55N>  
 PEND100: <https://www.statistikbanken.dk/PEND100>  
 ERHV6: <https://www.statistikbanken.dk/ERHV6>  
 UVM (Normering grundskoler): <https://uddannelsesstatistik.dk/Pages/Reports/1577.aspx>  
 UVM (Normering gymnasier): <https://uddannelsesstatistik.dk/Pages/Reports/1851.aspx>  
 UVM (Normering erhvervsuddannelse): <https://uddannelsesstatistik.dk/Pages/Reports/1850.aspx>  
 Daginistitutioner: <https://www.institutioner.dk/>



Figur 6. Flowdynamisk diagram af bevægelse gennem sygdomsstadier.

Agenter i modellen kan være i et af følgende sygdomsstadier: Modtagelig (S), Eksponeret (E), Præ-symptomatisk (IPS), Symptomatisk (I), Asymptomatisk (A), Rask (R) eller Død (D). Agenter, som befinder sig i det præ-symptomatiske, symptomatiske eller asymptomatiske stadiet, er infektions og kan således videreført smitte til agenter, som befinder sig i det modtagelige stadiet. Bliver en modtagelig agent inficeret, overgår de til at være eksponeret. Dette sygdomsstadiet repræsenterer den latente periode, hvor den inficerede agent endnu ikke er infektios. Agenterne kan bevæge sig gennem sygdomsstadierne, som vist på det flowdynamiske diagram, figur 6. Modellen antager, at 2/3 af agenterne bliver symptomatiske og at 1/3 forbliver asymptomatiske ved infektions tilstand. En andel symptomatiske agenter får et behandlingsbehov i løbet af deres sygdomsforløb og bliver indlagt på et Hospital (H). Sandsynligheden for indlæggelse blandt symptomatiske agenter er opdelt efter regioner og 10-års aldersgrupper baseret på data over indlæggelser i Danmark i september-oktober 2020.

Når en agent skifter til et nyt sygdomsstadiet, tildeles de den ventetid, som de skal opholde sig i stadiet. Ventetiden i de forskellige stadier er beskrevet ved gamma-fordelinger med parametre, som vist i tabel 3. Modellen simuleres i diskret tid. Hvert tids-skridt svarer til en halv dag.

Tabel 3. Parametre og quartiler for varighed af de enkelte stadier.

Stadier	Parametre		Kvartiler			Referencer
	Shape	Periode [Dage]	Nedre kvartil [Dage]	Median [Dage]	Øvre kvartil [Dage]	
Eksponeret (E)	3	3	2	3	4	Litteratur se referenceliste i bilag 5
Præsymptomatisk (IPS)	5	1,25	1	2	2	Litteratur se referenceliste i bilag 5
Symptomatisk (I)	4	7	5	7	9	Litteratur se referenceliste i bilag 5
Asymptomatisk (A)	4	7	5	7	9	Litteratur se referenceliste i bilag 5



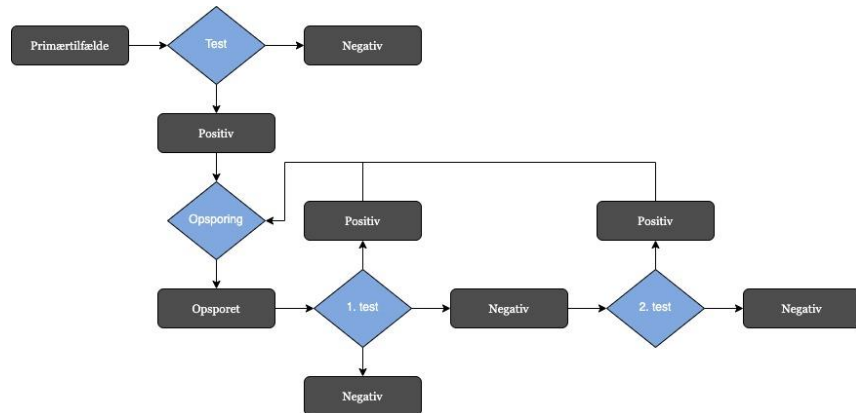
Hospitaliseret (H) Under 60 år	2	3	2	3	5	Linelisten SSI
Hospitaliseret (H) 60 år og der- over	2	5	3	5	7	Linelisten SSI
<b>Ventetider</b>						
timeSymp- ToOrderTest	5	1	1	1	1	Antagelser - af- venter STPS data
timeOrderTo- Test	2	2	1	2	3	Antagelser - af- venter STPS data
timeTestToRe- sult	6	1,5	2	2	2	Ventetider fra samfundssporet
traceDelay	5	1	1	2	3	Antagelser – af- venter STPS data

Sandsynligheden for, at en modtagelig agent bliver inficeret af en infektiøs agent og overgår til at være eksponeret i et givent netværk stiger med antallet af infektiøse agenter i netværket, de infektiøse agenter i netværkets smitsomhed, samt antallet af kontakter som både de modtagelige og infektiøse agenter har i netværket. Raten hvormed en modtagelig agent bliver inficeret er summen af smitterater fra de enkelte netværk, som agenten deltager i. Test og opsporing er indført i modellen ved følgende regler:

- Når en agent får symptomer, er der en sandsynlighed ( $pTestGivenSymptoms = 80\%$ ) for, at de bestiller en test efter en gammafordelt ventetid (timeSympToOrderTest). Hvis der er bestilt en test, vil personen reducere sine kontakter til 50% (undtagen i husholdninger, hvor kontakter reduceres til 70%).
- Der er en gammafordelt ventetid fra testen bestilles, til testen udføres (timeOrderToTest).
- Der er en gammafordelt ventetid fra testen udføres, til der kommer svar (timeTestToResult).
- Hvis der kommer positivt svar, vil agenten isolere sig yderligere; kontakter reduceres til 10% (husholdning: 50%). Derudover påbegyndes opsporing af netværk under følgende regler:
  - I skoleklasser, ungdomsskoleklasser, institutioner og i husholdninger opspores alle personer (i husholdninger foregår det dobbelt så hurtigt som i de øvrige netværk).
  - På kontorer (arbejdspladser) og i omgangskredse opspores et antal nære kontakter givet ved fordeling af kontakter under 1m i data fra HOPE projektet.
  - Personer, som tidligere er testet positiv, får ikke tildelt yderligere test.
  - Der opspores med en gammafordelt forsinkelse (traceDelay) fra den positive test.
  - Ved opsporing efter en person testes positiv tildeles de opsporedes personer testtider relativt til 48 timer før den positive fik symptomer - eller blev testet i et asymptomatisk tilfælde. Hvis muligt, gives test på dag 4 og dag 6, ellers dag 5 og 7, og ellers én test hurtigt muligt.
  - Personer, som er i et igangværende opsporingsforløb, får kun tildelt test, hvis de venter på mindre end to testsvar.
  - Den opsporedes person har samme ventetider på testsvar, som symptomatiske personer.



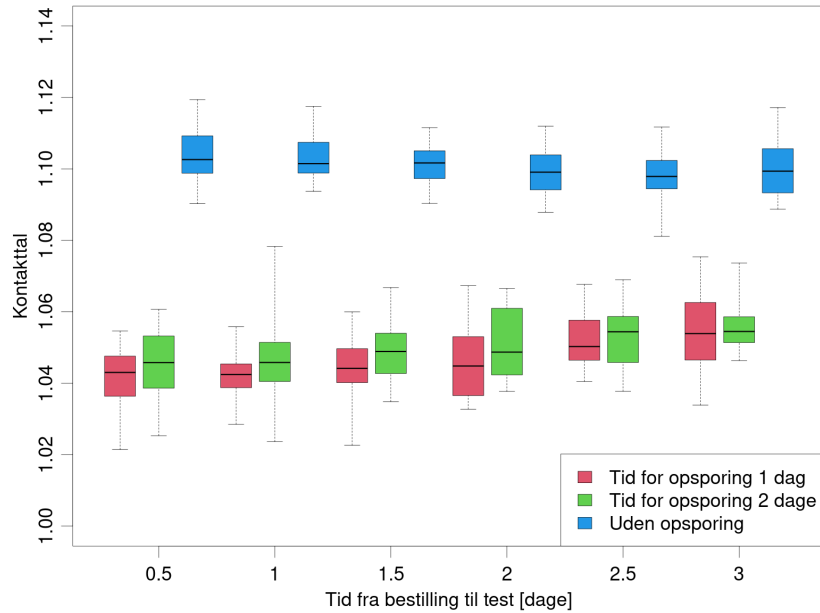
- Mens der ventes på test og testsvar, isoleres den opspored person på samme måde som en symptomatisk, der venter på svar.
- Hvis en opsporet person får negativt svar på den første test, vil der være en sandsynlighed for ( $pNoShow2ndTest = 40\%$ ) at de ikke tager test nummer 2.
- Efter et negativt svar på test nummer 1, vil isolationen brydes. Hvis der fås et positivt svar, inden test nummer 2 er taget, annuleres test nummer 2, og personens egne netværk opspores.
- For alle tests – om det er en opsporet person eller ej – antages der en sandsynlighed på 20% for en falsk negativ test (Kucirka et al., 2020).



Figur 7. Diagram, der viser test og opsporing i den agentbaserede model fra DTU.

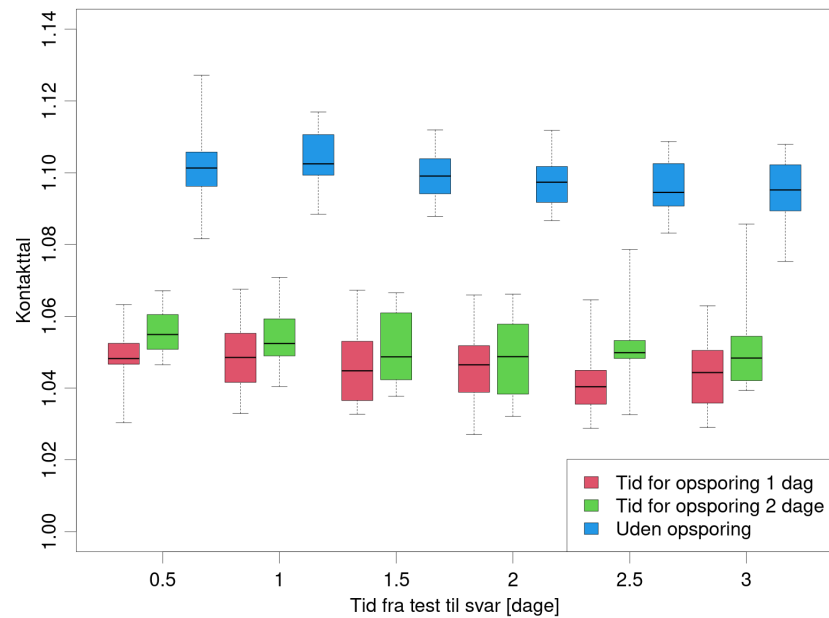


### Yderligere resultater



Figur 8. Kontakttallets afhængighed af ventetiden på at få taget en test hos primærtildældet, samt betydningen af hvor lang tid der går før nære kontakter opspores og går i tilsvarende isolation. For hver parameterværdi er der foretaget 40 simulationer og boxplottet viser median, de indre kvartiler samt minimum og maksimum af disse.

Af figur 8 fremgår det, at der ikke er nævneværdig forskel på reduktionen i kontakttallet, hvorvidt man reducerer ventetiden til at primærtildældet testes, i forhold til at reducere ventetiden til opsporing af nære kontakter.



Figur 9. Kontakttallets afhængighed af tiden fra der testes fra der foreligger et testsvar, samt betydningen af hvor lang tid der går indtil nære kontakter opspores og går i tilsvarende isolations. For hver parameterværdi er der foretaget 40 simulationer og boxplottet viser median, de indre kvartiler samt minimum og maksimum af disse.

Af figur 9 fremgår det, at der ikke er nævneværdig forskel på reduktionen i kontakttallet, hvorvidt man reducerer ventetiden fra at primærtifeldet og opspored kontakter testes til der foreligger et testsvar, i forhold til at reducere ventetiden til opsporing af nære kontakter. En årsag kan være, at ventetiden til testsvar gør, at en masse opspored og modtagelige kontakter er isoleret i længere tid og derfor ikke bliver smittet. Det er ikke undersøgt om dette kun ses når kontakttallet er nær 1.



## Bilag 3. Regneeksempel

Følgende er et illustrativt regneeksempel på den agentbaserede model fra Niels Bohr Institutet beskrevet i bilag 1. Udregningerne er baseret på modellens underliggende antagelser, nemlig at perioden for eksposition (E ( $T_E$ )), hvor den latente fase er en gammafordeling med middelværdi på 4.7 dage, og perioden for den smitsomme fase er en gammafordeling med middelværdi på 7 dage, samt en antagelse om, at 40% af cases findes uden kontaktopsporing. Det antages, at for de COVID-19 tilfælde der findes uafhængigt af kontaktopsporingen, er de smittet uniformt fordelt i den smitsomme periode (I). Vi udregner nu tiden man er asymptomatisk men smitsom ved at trække tal fra fordelingen af tider for hele perioden, man er smitsom og tester en andel p, på et uniformt tilfældigt tidspunkt. Det giver en fordeling og en gennemsnitlig eksponeringstid (se figur 10A).

Vi kigger nu på et sekundært tilfælde, der blev smittet på et uniformt tilfældigt tidspunkt i den smitsomme periode for primærtildældet. Denne person kan enten findes tilfældigt, eller ved at primærtildældet testes, og at sekundærtildældet opspores efter en tidsperiode (d for delay). Denne ventetid, er tiden fra at primærtildældet testes til at sekundærtildældet kontaktes, og afspejler således både ventetid til test samt ventetid til opsporing. Igennem antages det, at sekundærtildældet går i isolation øjeblikkeligt. Ved igen at trække tal tilfældigt fra de relevante fordelinger fås en eksponeringsperiode, hvori sekundærtildældet måske opspores, forhåbentligt inden smitten er ført videre.

### Resultat

I figur 10B vises det gennemsnitlige antal dage en kontakt er eksponeret for smitte, som en funktion af den samlede ventetid til test og opsporing. Herudfra estimeres effekten af kontaktopsporing på det effektive kontakttal, Rt. Det antages, at en given andel (fc) af alle smittetildældet, findes via kontaktopsporing, og derved reduceres smitten, idet eksponeringsperioden for opspored kontakter reduceres. Herved fås et simpelt estimat af effekten af kontaktopsporing på kontakttallet Rt. Dette vises i figur 10C. Farverne på graferne viser, hvor stor en andel af smitten der kan reduceres, såfremt eksponeringsperioden reduceres, som følge af kontaktopsporing. Hvis det f.eks. antages, at der er 2000 nye smittede med COVID-19 per dag (ca. 1000 fundne smittede + et mørketal), så svarer 0.05 grafen (orange) til at 100 smittede bliver fundet gennem kontaktopsporing dagligt.

En væsentlig begrænsning er, at disse udregninger ikke medtager effekten af, at flere COVID-19 tilfælde bliver fundet pga. kontaktopsporing, men er udelukkende baseret på effekten ved at forlænge eksponeringsperioden for kontakter.

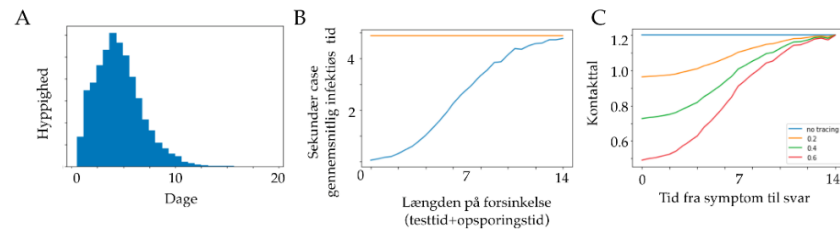
I modellen indgår 4 mulige eksponeringsperioder. 1: Kontakter opspores ikke, hvorved eksponeringsperioden ikke afkortes (blå graf), 2: 20% af kontakter opspores (gul graf), 3: 40% af kontakter opspores (grøn graf) og 4: hvis 80% af kontakter opspores (rød graf).

Af regneeksemplet fremgår det, at givet antagelserne i eksemplet vil kontakttallet kunne reduceres med ca. 50%, såfremt man opsporer 50% af alle kontakter inden for ca. 3 dage.

Bemærk at alle kurverne i figur 10C er meget flade i intervallet mellem dag 0 og 3. Dette betyder, at der kun opnås en lille gevinst ved at afkorte den samlede ventetid fra symptomer til der foreligger et testsvar inden for denne periode, men at der til gengæld er en stor gevinst ved at øge andelen af opspored kontakter.



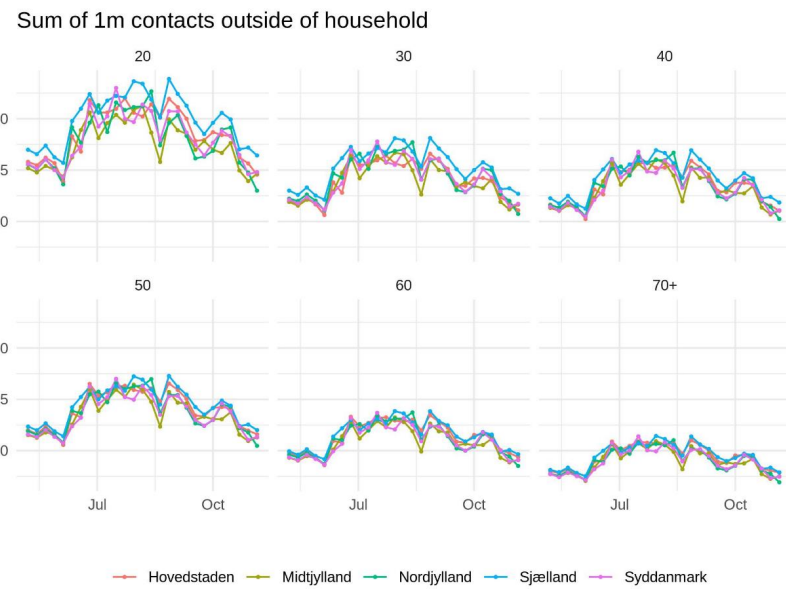
Det skal i øvrigt bemærkes, at det i eksemplet antages, at opsporede kontakter går i isolations, indtil de får svar på deres test.



*Figur 10:A) Fordeling af eksponeringstiden, gennemsnit = 4.9 dage. B) Gennemsnitlig eksponeringstid for sekundære tilfælde (blå), som funktion af den samlede ventetid til test og opsporing. Den orange graf viser gennemsnittet i ventetiden til test og opsporing for primært tilfældet som reference. C) Det effektive kontakttal  $R_t$  efter kontaktsporing som funktion af ventetiden fra symptomer til testsvar hvor udgangspunktet er et kontakttal på 1.2, inden der iværksættes opsporing. Farverne indikerer hvor stor en andel af kontakter der opspores, hvorved eksponeringstiden reduceres.*



## Bilag 4. Udvikling i antal kontakter fra HOPE projektet



*Figur 11. Kilde: Hope-projektet (12.11.2020). Estimating Local Protective Behavior in Denmark with dynamic MRP. [https://github.com/mariefly/HOPE/raw/master/HOPE\\_report\\_2020-11-12.pdf](https://github.com/mariefly/HOPE/raw/master/HOPE_report_2020-11-12.pdf)*



## Bilag 5. Beskrivelse af parametre brugt i rapporten

Modellerne i rapporten bygger på en række parametre. Estimaterne, som parametre er baseret på er udvalgt af den relevante institution, der har udarbejdet modellerne. Begrundelsen for valg af estimaterne er beskrevet nærmere i dette bilag.

Overordnet set er parametre om sygdomsforløb primært baseret på international litteratur på emnet, men også på data fra den danske befolkning. Estimater over befolkningens adfærd i forbindelse med covid-19 bygger på en række danske undersøgelser fra i år, samt på data over danskernes rejsemønstre.

### Estimater for latensperiode, inkubationsperiode og infektions periode fra litteraturen:

Særligt relevant for simuleringerne over effekten af kontaktopsporing er estimaterne bag sygdomsforløbet, herunder hvor lang tid der går fra eksponering til, at vedkommende kan smitte, og derefter til, at vedkommende vises symptomer. Estimaterne i modellen er blandt andet baseret på andre forskeres data, som er offentliggjort i international litteratur om covid-19.

For at finde de bedste estimat på *latensperioden* har modelgruppen trianguleret distributioner fra nedenstående kilder. Estimatet er 3,6 dage med et interval på mellem 3-5 dage.

- Read et al. (2020). Novel coronavirus 2019-nCoV: Early estimation of epidemiological parameters and epidemic predictions. *Preprint*.
- Li et al. (2020). Early Transmission Dynamics in Wuhan, China, of Novel Coronavirus–Infected Pneumonia. *N. Engl. J. Med.*
- Li et al. (2020). Substantial undocumented infection facilitates the rapid dissemination of novel coronavirus (SARS-CoV2). *Science*.
- Milne and Xie (2020). The Effectiveness of Social Distancing in Mitigating COVID-19 Spread: a modelling analysis. *Preprint*.

For at finde det bedste estimat af *inkubationsperioden*, har Ekspertgruppen gennemgået nedenstående litteratur. Estimatet er 5 dage med et interval på mellem 3-7 dage.

- Lauer et al. (2020). The Incubation Period of Coronavirus Disease 2019 (COVID-19) From Publicly Reported Confirmed Cases: Estimation and Application. *Ann. Int. Med.*
- Li et al. (2020). Early Transmission Dynamics in Wuhan, China, of Novel Coronavirus–Infected Pneumonia. *N. Engl. J. Med.*
- Anderson et al. (2020). How will country-based mitigation measures influence the course of the COVID-19 epidemic. *The Lancet*.
- Linton et al. (2020). Incubation Period and Other Epidemiological Characteristics of 2019 Novel Coronavirus Infections with Right Truncation: A Statistical Analysis of Publicly Available Case Data. *J. Clin. Med.*
- Liu et al. (2020). Transmission dynamics of 2019 novel coronavirus (2019-nCoV). *bioRxiv*.
- Shen et al. (2020). Modelling the epidemic trend of the 2019 novel coronavirus outbreak in China. *bioRxiv*.



- Backer et al. (2020). Incubation period of 2019 novel coronavirus (2019-nCoV) infections among travellers from Wuhan, China, 20–28 January 2020. *Euro Surveill.*
- Gostic et al. (2020). Estimated effectiveness of symptom and risk screening to prevent the spread of COVID-19. *eLife*
- Hellewell et al. (2020). Feasibility of controlling COVID-19 outbreaks by isolation of cases and contacts. *The Lancet Global Health.*
- Milne and Xie (2020). The Effectiveness of Social Distancing in Mitigating COVID-19 Spread: a modelling analysis. *Preprint.*

For estimatet af *den infektiose periode*, hvor det bedste estimat er 5 dage, mens det bedste interval er mellem 3-7 dage, har Ekspertgruppen gennemgået følgende artikler:

- Read et al. (2020). Novel coronavirus 2019-nCoV: Early estimation of epidemiological parameters and epidemic predictions. *Preprint.*
- Prem et al (2020). The effect of control strategies that reduce social mixing on outcomes of the COVID-19 epidemic in Wuhan, China. *Centre for the Mathematical Modelling of Infectious Diseases COVID-19 Working and Jit, Mark and Klepac, Petra, The Effect of Control Strategies that Reduce Social Mixing on Outcomes of the COVID-19 Epidemic in Wuhan, China.*
- Milne and Xie (2020). The Effectiveness of Social Distancing in Mitigating COVID-19 Spread: a modelling analysis. *Preprint.*

#### **HOPE rapporter og data:**

En del af estimererne i modellerne om befolkningens adfærd, herunder kontaktmønstre, bygger på både data og rapporter for Hope-projektet (<https://hope-project.dk/#/>).

HOPE-projektet udsender løbende spørgeskemaer til tilfældigt udvalgte personer i Danmark vedrørende både deres tillid til myndighederne, og til deres adfærdsmønstre, herunder hvor mange de ses med i forskellige kontaktkategorier, hvor meget afstand de holder fra andre mennesker etc. Denne information samles i rapporter, der løbende offentliggøres.

Udover HOPE-rapporten, der henvises til i Bilag 4 ([https://github.com/mariefly/HOPE/raw/master/HOPE\\_report\\_2020-11-12.pdf](https://github.com/mariefly/HOPE/raw/master/HOPE_report_2020-11-12.pdf)), oversender HOPE-projektet løbende anonymiserede data om befolkningens adfærd under covid-19 til Ekspertgruppen, der anvender det i deres modeller. Ekspertgruppen har også adgang til HOPE-projektets rapporter, der sammenskriver data.

#### **Trafik data:**

Antagelser om befolkningens adfærd bygges ligeledes på trafikdata, hvorudfra man kan bestemme danskernes rejsemønstre. Efter aftale med Trafik-, Bygge- og Boligstyrelsen får Ekspertgruppen løbende adgang anonymiserede data over danskernes bevægelse rundt i landet. Data er bl.a. brugt til at bestemme den typiske afstand mellem kontakter og afstanden mellem afstands-uafhængige kontakter. Data bygger på 5 forskellige kilder:

- Overblik over rejsende, der bruger rejsekort, som kommer fra Rejsekort og Rejseplanen A/S
- Overblik over biltrafik på Øresunds- og Storebæltsbroen fra Sund og Bælt A/S



- Overblik over flytrafik (antal passagerer) til og fra Københavns Lufthavn og Billund Lufthavn
- Overblik over biltrafikken på Statsvejsnettet og cykeltrafikken (samlet ud fra tællestandere) leveret af Vejdirektoratet.
- Overblik og færgetrafik på 5 rederier, der dækker over 17 færgeruter. Data er leveret af Danske Rederier.

#### **Estimater for ventetider til test**

Estimater for ventetider til test og svar på test er taget fra TCDKs hjemmeside (<https://tcdk.ssi.dk/vente-og-svartider>).

#### **Data fra SSIs Linelisten**

Linelisten på SSI indeholder informationer om de covid-19 podninger, der tages en given dag. Data fra Linelisten er bl.a. brugt til at modellere risikoen for at blive hospitaliseret i løbet af et covid-19-forløb for personer over og under 60 år.

#### **Spørgeskemaundersøgelse blandt covid-19 syge lavet af SSI i foråret:**

I foråret 2020 foretog SSI en telefonisk spørgeskemaundersøgelse blandt en række personer, der fik konstateret covid-19. Spørgsmålene undersøgte deltagernes sygdomsforløb, herunder symptomer, hvorvidt nære kontakter i husstanden var smittet og lignende.

Data fra spørgeskemaundersøgelsen blev i modellerne brugt til at estimere tiden fra symptomdebut til tests i dage.

#### **Den nationale prævalensundersøgelse for covid-19:**

SSI iværksatte i maj en undersøgelse af, hvor udbredt covid-19 var blandt danskerne. Undersøgelsen bestemmer seroprævalencen blandt et repræsentativt udsnit af danskerne fra maj og til i dag. Informationer fra prævalensundersøgelsen har været anvendt i modellerne til at estimere sandsynligheden for at få symptomer og blive testet.



## Bilag 6. Medlemmer af ekspertgruppen

Ekspertgruppen ledes af læge Camilla Holten Møller og overlæge Robert Leo Skov, Infektionsberedskabet, Statens Serum Institut.

### **Danmarks Tekniske Universitet, Institut for Matematik og Computer Science**

- Kaare Græsbøll, ph.d., MSc, Seniorforsker, Sektion for dynamiske systemer
- Lasse Engbo Christiansen, ph.d., MSc Eng, lektor, Sektion for dynamiske systemer
- Sune Lehmann, Professor, Afdelingen for Kognitive Systemer
- Uffe Høgsbro Thygesen, Civilingeniør, ph.d., lektor, Sektion for dynamiske systemer

### **Københavns Universitet, Det Sundhedsvidenskabelige Fakultet, Institut for Veterinær- og Husdyrvidenskab,**

- Carsten Thure Kirkeby, Seniorforsker, ph.d., MSc. Sektion for Animal Welfare and Disease Control
- Matt Denwood, BVMS, ph.d., Sektion for Animal Welfare and Disease Control

### **Københavns Universitet, Institut for Folkesundhedsvidenskab**

- Theis Lange, Vice Instituteder, Lektor i Biostatistik, ph.d., Biostatistisk Afdeling

### **Københavns Universitet, Niels Bohr Institutet**

- Troels Christian Petersen, Lektor, Eksperimentel subatomar fysik

### **Roskilde Universitets Center, Institut for Naturvidenskab og Miljø**

- Viggo Andreasen, Lektor, Matematik og Fysik

### **Region Hovedstaden**

- Anders Perner, Professor, Overlæge, Intensivafdelingen, Rigshospitalet

### **Danmarks Statistik**

- Laust Hvas Mortensen, Chefkonsulent, professor, ph.d., Metode og Analyse

### **Statens Serum Institut**

- Mathias Heltberg, Postdoc ENS Paris samt Statens Serum Institut. Infektionsberedskabet
- Frederik Plesner Lyngse, Postdoc, Økonomisk Institut, Københavns Universitet samt Statens Serum Institut, Infektionsberedskabet
- Peter Michael Bager, Seniorforsker, ph.d., Infektionsberedskabet, Epidemiologisk Forskning, Statens Serum Institut
- Robert Skov, Overlæge, Infektionsberedskabet, Statens Serum Institut
- Camilla Holten Møller, Læge, PhD, Infektionsberedskabet, Statens Serum Institut



## D *SSI Notat*

The following pages contain the report from Statens Serum Institut, the Danish CDC:

Ekspertgruppen for matematisk modellering, “*Scenarier for udviklingen i den engelske virusvariant af SARS-COV-2 (cluster B.1.1.7)*” (Statens Serum Institut, 2021).

The report is from January 2, 2021 and is a summary of the estimated spread of the “alpha” variant of COVID-19 (B.1.1.7) in Denmark. The report is in Danish and is based on two models, one from DTU and our agent based model from NBI.



d. 2. januar 2021

*Notatet er opdateret d. 22. januar 2021 med en præcisering af formuleringer vedrørende udviklingen i forholdet mellem Cluster B.1.1.7 og øvrige virusvarianter.*

#### **Scenarier for udviklingen i den engelske virusvariant af SARS-COV-2 (cluster B.1.1.7)**

Ekspertgruppen for matematisk modellering, der ledes fra SSI, bringer i dette notat en række estimer for den forventede udbredelse af cluster B.1.1.7 i den kommende periode, dels ved logistisk regression af udviklingen i forekomsten af varianten, og dels ud fra simuleringer af spredningen af varianten i en agentbaseret model.

#### **Sammenfatning**

- Den observerede udvikling i forekomsten af cluster B.1.1.7 i Danmark, svarer til en ugentlig vækstrate for forholdet mellem cluster B.1.1.7 og de øvrige virusvarianter på 72% (95% CI: [37, 115] %).
- Med udgangspunkt i den aktuelle situation hvor 2,3% af virusvarianterne i den rutinemæssige helgenomsekventering tilhører cluster B.1.1.7, estimeres det, at varianten vil udgøre halvdelen af de cirkulerende virusstammer i Danmark om 40-50 dage såfremt ovennævnte stigning fortsætter.
- Det nuværende niveau af restriktioner forventes ikke at være tilstrækkeligt til at få kontakttallet for cluster B.1.1.7 under 1. Derfor vil denne vokse eksponentielt upåagtet at det samlede kontakttal (for alle virusvarianter) kan være under 1 indtil cluster B.1.1.7 overtager om omkring en måned.
- Forekomsten af cluster B.1.1.7 er højest i Region Nordjylland, og udviklingen i forekomsten er ca. fire uger foran Region Hovedstaden.
- Det er på baggrund af engelske data estimeret at kontakttallet er ca. 1,5 gange højere for den nye virusvariant i forhold til andre virusvarianter.
- Den reduktion i smittetal og indlæggelser, der kan opnås i den kommende måned vil give et lavere udgangspunkt for den forøgede smitte og stigende kontakttal, som vi må forvente.

Disse beregninger er behæftet med usikkerheder af forskellige grunde. I perioden op til jul var der stor efterspørgsel på tryghedstest, og i samme periode er der udført et stigende antal antigen test. Derimod så vi i juledagene, at kun ganske få har ladet sig teste. Disse ændringer i testdynamikker gør det svært at følge udviklingen i covid-19, idet de vanlige indikatorer såsom incidenser, positivprocenter og kontakttallet påvirkes af den ændrede fordeling af covid-19-positive blandt de testede. Et lignende mønster forventes i dagene op til og efter nytår. Desuden har vi endnu ikke set effekten af de sidst indførte tiltag, herunder lukning af detailhandlen og liberale erhverv. Samlet set giver dette usikkerhed omkring det aktuelle kontakttal. Analysen er baseret på 76 isolater med cluster B.1.1.7 fordelt på de fem regioner. Den lille stikprøve giver relativt store statistiske usikkerheder. Der vil derfor være behov for at løbende at opdatere estimaterne og lave nye analyser.



### Logistisk regression for spredningen af cluster B.1.1.7

Som det fremgår af nedenstående tabel, er der stor forskel på, hvornår man har fundet cluster B.1.1.7 i de enkelte regioner.

*Tabel 1. Forekomst af cluster B.1.1.7 i de fem regioner baseret på helgenomsekventering af stikprøver af SARS-CoV-2 positive isolater.*

Uge	Hovedstaden		Midtjylland		Nordjylland		Sjælland		Syddanmark	
	B.1.1.7	Total	B.1.1.7	Total	B.1.1.7	Total	B.1.1.7	Total	B.1.1.7	Total
45	0	656	0	283	0	238	0	181	0	200
46	4	420	0	327	0	305	0	132	0	168
47	0	588	0	297	0	240	0	143	0	241
48	3	679	0	291	0	169	0	165	0	195
49	0	825	0	332	3	64	0	246	0	208
50	2	892	0	360	7	92	0	214	1	431
51	3	753	0	524	9	254	3	310	4	354
52	8	774	5	221	12	169	10	193	1	225

Ud fra udbredelsen af cluster B.1.1.7 i Danmark samt andelen af nye isolater i overvågningen som er relateret til clusteret, anvendes logistisk regression til at estimere den forventede udbredelse af cluster B.1.1.7. Da fokus er på spredningen af virusvarianten, og ikke på introduktioner af denne, er det kun regioner, hvor der er detekteret isolater tilhørende cluster B.1.1.7 i mindst fire uger – dvs. Region Hovedstaden og Region Nordjylland, der er medtaget i denne første analyse.

Der er lavet logistisk regression med uge og region som forklarende variable. Der er også testet en interaktion, men den er ikke signifikant.

*Tabel 2. Estimater for logistisk regression af andelen af cluster B.1.1.7. Referencen repræsenterer Region Hovedstaden.*

	Estimate	Std. Error	z value	Pr{> z }
(Intercept)	-32.812	5.679	-5.778	0.000
Uge	0.540	0.112	4.844	0.000
Region Nordjylland	2.221	0.311	7.133	0.000



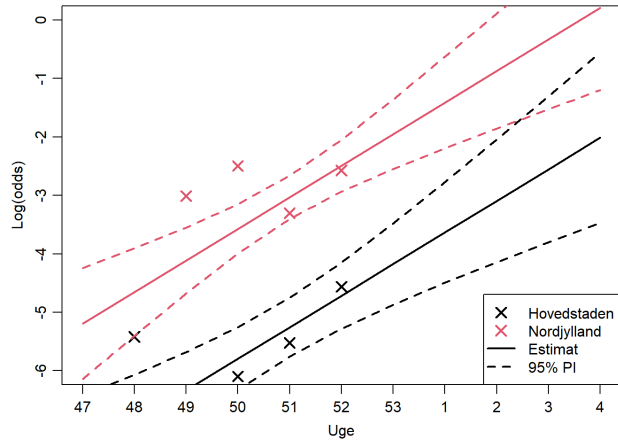
Det ses, at log(odds) for at detektere cluster B.1.1.7 er 2.2 højere i Region Nordjylland end i Region Hovedstaden. Det svarer til odds på 9.2. Det mest interessante er den tidslige udvikling, hvoraf det ses at log(odds) øges med 0.54 for hver uge. Dette svarer til at cluster B.1.1.7 har en ugentlig vækstrate i odds (forholdet mellem antal cluster B.1.1.7 og øvrige virusvariante) på 72% (95% CI: [37, 115] %), hvilket med den nuværende lave andel af cluster B.1.1.7 svarer til den samme stigning i andelen af cluster B.1.1.7 blandt alle positive prøver. Usikkerheden på estimatet er endnu ganske stort og estimatet er følsomt over for hvilke uger der medtages. Uanset usikkerheder, svarer det fundne estimat til de der er rapporteret fra England for denne virusvariant og det tyder på, at cluster B.1.1.7 har samme forøgede transmissionsrate i Danmark som i England.

Det ses, at log(odds) for at detektere cluster B.1.1.7 er 2.2 højere i Region Nordjylland end i Region Hovedstaden. Det svarer til odds på 9,2, dvs. at sandsynligheden for at detektere cluster B.1.1.7 her er 9,2 gange højere. Det svarer også til at Region Nordjylland er fire uger foran Region Hovedstaden i andelen af cluster B.1.1.7.

Det forventes, at usikkerhederne vil blive reduceret væsentligt når der er data for 1-2 uger mere. Men givet at B.1.1.7 er så meget mere smitsom end hidtidige varianter vil det kræve længerevarende restriktioner at sænke smittetallet.

De seneste estimerer af kontakttallet er lige under 1,0. Dette er dog påvirket af den ændrede testaktivitet og adfærd hen over jul og nytår, og vi har endnu ikke et overblik over konsekvenserne af sammenkomster i forbindelse med jul og nytår. Endvidere har vi endnu ikke set effekten af nedlukningen af de liberale erhverv og detailhandlen omkring jul. Derfor er det forventningen, at en fastholdelse af de nuværende restriktioner vil give et fald i kontakttallet, hvis man kigger på de virusvariante som vi har set for introduktionen af cluster B.1.1.7. I England har man estimeret, at deres reference kontakttal var 0,8 for andre virusvariante og 1,2 for cluster B.1.1.7. Det observerede kontakttal er et vægtet gennemsnit af virusvarianterne i populationen.

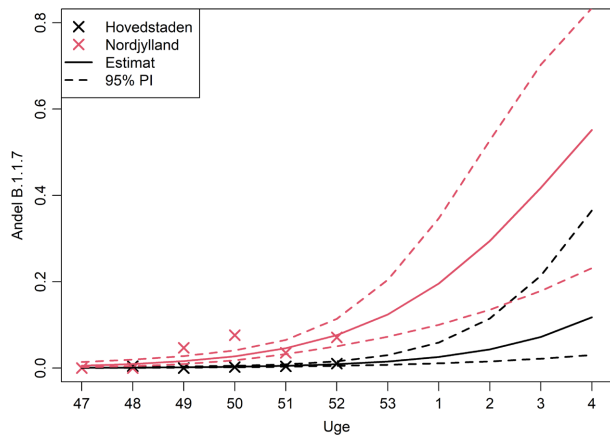
Figur 1 viser en fremskrivning af log(odds) for B.1.1.7 mod andre virusvariante baseret på ovenstående logistiske regression. Estimatet er, at cluster B.1.1.7 allerede i uge 4 vil udgøre halvdelen af alle positive test i Region Nordjylland. Dette er dog behæftet med stor usikkerhed på baggrund af de nuværende data.



Figur 1. Log(odds) for at detektere cluster B.1.1.7 i hhv. Region Hovedstaden og Region Nordjylland

Ved sammenligning med England er vi nu, hvor de var i starten af november, hvor South East havde log(odds) på -2 svarende til Nordjylland og både London og East of England havde log(odds) omkring -4 svarende til Hovedstaden<sup>1</sup>

Figur 2 viser den samme fremskrivning som i figur 1. Blot er der transformerede tilbage til andelen af positive test, som tilhører cluster B.1.1.7.



<sup>1</sup> 2020\_12\_23\_Transmissibility\_and\_severity\_of\_VOC\_202012\_01\_in\_England.pdf  
(cmmid.github.io)



*Figur 2. Udviklingen i forekomsten af cluster B.1.1.7 i de kommende uger. Fremskrivningen viser, at halvdelen af isolaterne i Region Nordjylland vil være cluster B.1.1.7 omkring uge 4.*

Det skal bemærkes, at udviklingen i Hovedstaden er ca. 4 uger efter udviklingen i Nordjylland. Det er endnu for tidligt at udtales sig om niveauet i de andre tre regioner, men særlig Region Sjælland synes at have oplevet en hurtig stigning, om end det er baseret på meget lidt data. De næste par uger vil forbedre estimatet af niveauet i alle regioner.

Hen over julen har der været et nyt toppunkt i antal indlagte og der er endnu kun set små fald. Det er først i uge 1, at vi kan forvente at se eventuelle indlæggelser som følge af smitte i julen. Alt andet lige må dette forventes at give en yderligere kortvarig pukkel i antal nye indlæggelser.

På nuværende tidspunkt er prognosen, at vi har omkring en måned før det samlede kontakttal for alle virusvarianter hurtigt vil stige på grund af øget udbredelse af cluster B.1.1.7. Hvis restriktionerne skærpes i den kommende tid, vil det give en reduktion i smittetal og indlæggelser og dermed et lavere udgangspunkt for den forøgede smitte og stigende kontakttal, som vi må forvente.

Et første estimat af kontakttallet for cluster B.1.1.7 for perioden uge 47 til 52 og baseret på observationer fra Region Hovedstaden og Region Nordjylland er 1.5 (95% CI [1,2 ; 1,7]) - dette er estimeret vha. Poisson regression med offset lig med  $0.7 * \log(\text{antal sekventerede})$ . Det gennemsnitlige kontakttal (baseret på SSIs publicerede kontakttal 2020-12-29) for perioden er 1,1. Da kontakttallet for cluster B.1.1.7 er så meget højere må det selv med de nuværende restriktioner forventes, at det vedbliver med at være over 1 og dermed forventes cluster B.1.1.7 at vokse eksponentielt, hvis det nuværende niveau af restriktioner fastholdes.

#### **Simulering af spredningen af cluster B.1.1.7 i en agentbaseret model**

##### *Agentbaserede modeller*

Spredningen af cluster B.1.1.7 er simuleret i en agentbaseret model, som er udviklet af Niels Bohr Institutet, Københavns Universitet (NBI). En agentbaseret model simulerer et antal agenter (individer i en population) og deres interaktioner med andre agenter, svarende til de interaktioner som en befolkning normalt viser. Hver agent repræsenterer således en person, som er knyttet til en lokation i Danmark, svarende til deres bopæl. Agenterne indgår i flere forskellige netværk, f.eks. husstand, job og skole hvor de har kontakt til andre personer. Derudover har de kontakt til tilfældige personer i samfundet i den tid, hvor personen ikke er hjemme, på job eller i skole. Hvis en agent bliver smittet med SARS-CoV-2, er forløbet for den enkelte agent beskrevet således, at agenten først er eksponeret (E) og derefter infektiøs (I), hvorefter agenten ikke længere er smitsom og betragtes som rask (R). De gennemsnitlige tider i hvert sygdomsstadiu kan findes i bilag 1. Hver kontakt, som en agent eksponeres for, tildeles en sandsynlighed for at blive smittet af en anden agent, såfremt denne er smitsom. For en detaljeret beskrivelse af den agentbaserede model, herunder de inkluderede parametre, henvises til bilag 1.

##### *Forbehold*



Mens en agentbaseret model kan medtage mere detaljerede dynamikker i en epidemi, så kræver en præcis simulation input fra data, som ofte ikke er tilgængelige eller forefindes, fx hvem en person mødes med i løbet af en dag. Derfor kan en sådan model have unøjagtigheder eller bygge på antagelser, som ikke er retvisende. Det er ikke muligt at kvantificere den nøjagtige størrelse eller effekt af disse potentielle fejlkilder. Da datagrundlaget for disse simuleringer er sparsomt, fordi vi endnu har få datapunkter for cluster B.1.1.7, vil resultatet være behæftet med væsentlig usikkerhed.

#### *Resultater*

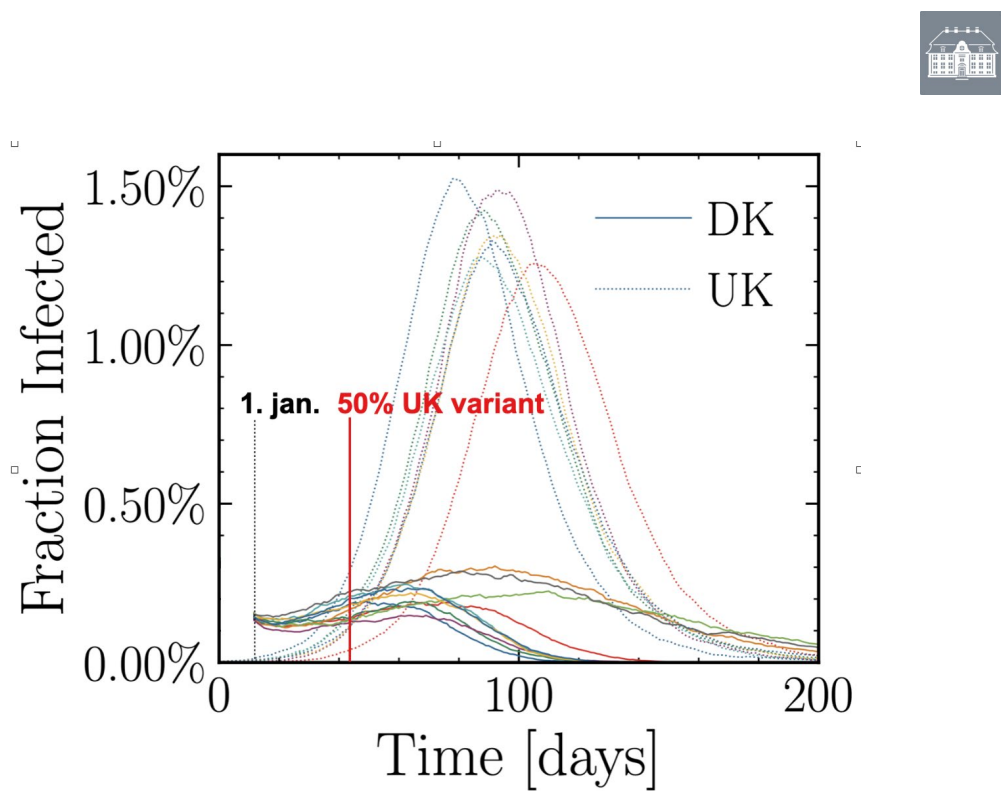
I det følgende er udviklingen simuleret i en model, hvor udgangspunktet er 1/10 af Danmarks befolkning, og hvor cluster B.1.1.7 fra starten udgør omkring 5% af de cirkulerende virusvarianter. Epidemien simuleres ud fra et kontakttal på omkring 1,0, samt en antagelse om, at cluster B.1.1.7 smitter 50% mere, som rapporteret fra England<sup>2</sup>

Figur 3 viser, hvordan en epidemi vil udvikle sig i tid, forudsat at det simulerede scenarie ikke ændres. Der opdeles i hhv. de nuværende virusvarianter (DK, fulde linjer) og det engelske cluster B.1.1.7 (UK, stiplede linjer). Simulationen er gentaget flere gange (forskellige farver) for at se, hvor store variationer der forekommer. Som det kan ses, så udfases DK-versionen af smitten, mens UK-versionen B.1.1.7 giver ophav til en eksponentiel vækst, idet kontakttallet for denne er væsentligt over 1.

Af figuren fremgår det, at cluster B.1.1.7 ca. 35-40 dage fra simulationens start ("1. jan.") udgør omkring 50% af de cirkulerende virusvarianter. Da simulationen er startet med en større andel UK-varianter (5%) end det aktuelle landsgennemsnit (2.3%), så bliver estimatet 40-50 dage til at halvdelen af de sekventerede varianter tilhører cluster B.1.1.7. I de viste simulationer er de første smittet med cluster B.1.1.7 varianten placeret i Hovedstadsområdet. I andre scenarier, hvor cluster B.1.1.7 varianten i starten udvikler sig i et tyndere befolket område tager udviklingen lidt længere tid, op til 60 dage.

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<sup>2</sup>2020\_12\_23\_Transmissibility\_and\_severity\_of\_VOC\_202012\_01\_in\_England.pdf  
(cmmid.github.io)



Figur 3. Den forventede udvikling i cluster B.1.1.7 sammenholdt med udviklingen i øvrige virusvarianter, simuleret i en agentbaseret model. Ud fra simulationerne estimeres det, at B.1.1.7 varianten vil være dominerende efter 40-50 dage.

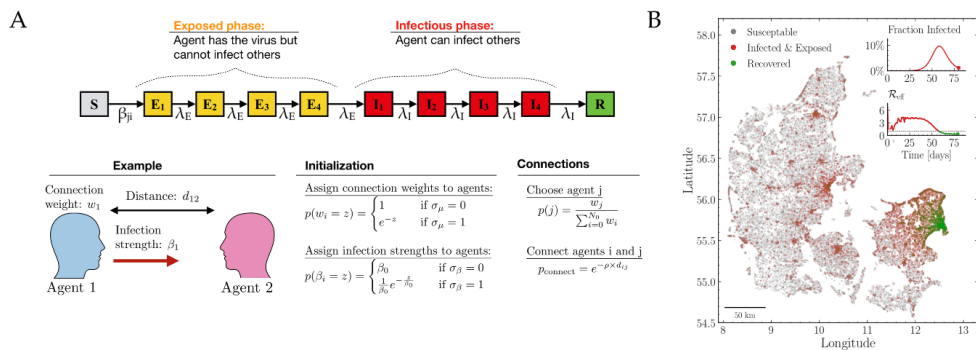


### Bilag 1. Beskrivelse af den agentbaserede model

Den nedenstående modelbeskrivelse er et uddrag fra ekspertrapporten "effekten af kontaktopsporing" der er publiceret d. 16. december 2020

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Den agentbaserede model baseres på agenter, dvs. individer hvis karakteristika er tildelt ud fra statistiske fordelinger i befolkningen. Dette er f.eks. en aldersfordeling og en fordeling over pendlerafstande. Modellen starter med at fordele Danmarks bopæle ud i landet baseret på det danske hussalg over de sidste 15 år. Herefter placeres agenter i hver husstand baseret på deres alder og geografiske placering.



Figur 5: A) Skematiske oversigt over hvordan interaktionsnetværket i modellen ser ud. B) Eksempel på simulation af smittespredning i Danmark i modellen, gennem et simuleret tilfælde af flokimmunitet i København.

Et afgørende element i modellen er opbygningen af alle personers interaktionsnetværk. Dette genereres ved, at hver agent har et netværk, de interagerer med. Dette opdeles i tre dele: 1) kontakter i hjemmet, 2) kontakter på arbejdet, 3) kontakter i kategorien andre kontakter. Der er ikke nogen geografisk afhængighed af antallet af kontakter på arbejdet, men i den kategori der kaldes "andre", vil der generelt være flere kontakter for dem der bor i tæt befolkede områder i forhold til dem der bor på landet. Måden hvorpå netværket dannes er vist i Figur 5A.



Ud fra data fra HOPE-projektet har vi estimeret, hvor mange personer hver agent vil interagere med, og i denne model vil alle agenter have mellem 3 og 15 daglige kontakter.

Når modellen simuleres vil alle inficerede agenter gennemgå et forløb, hvor de er i en latent periode, hvor de ikke smitter, hvorefter de vil rykke over i en infektiøs periode, hvor de kan smitte agenter i deres netværk. Denne model simuleres ud fra det der kaldes Gillespie algoritmen, således at netværket opdateres instantan for alle smittebegivenheder. En samling af de væsentligste parametre er vist herunder (Tabel 2).

*Tabel 2: Parametre i den agentbaserede model*

Parameter	Værdi interval for middelværdien	Reference
Antal kontakter per dag	3-15	HOPE projektet
Latent tid (dage)	3-5	Litteratur se referenceliste i bilag 5
Infektiøs tid (dage)	4-8	Litteratur se referenceliste i bilag 5
Andel af kontakter i ”andre” (%)	30-80	HOPE projektet
Typisk afstand mellem kontakter (km)	5-20	Trafik data
Andel afstandsufhængige kontakter (%)	3-5	Trafik data
Tid fra symptom til test (Dage)	0-2	Fordeling fra spørgeskemaundersøgelse i foråret 2020 (ikke offentliggjort)
Sandsynlighed for at få symptomer og blive testet (%)	20-60 %	Prævalensundersøgelsen
Sandsynlighed for at kontakte husstand (%)	100%	Antagelse
Sandsynlighed for at kontakte kollegaer (%)	40-80	Antagelse
Sandsynlighed for at kontakte andre (%)	0-75	Antagelse

This document was typeset using **LATEX** and modified version of the **tufte-style-thesis** class.  
The style is heavily inspired by the works of Edward R. Tufte and Robert Bringhurst.