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Automated Multi-Objective Calibration of Biological Agent-Based Simulations

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Abstract

Computational agent-based simulation is increasingly used to complement lab-15 oratory techniques in advancing our understanding of biological systems. Calibra-16 tion, the identification of parameter values that align simulation with biological 17 behaviours, becomes challenging as increasingly complex biological domains are 18 simulated. Complex domains cannot be characterised by single metrics alone, ren-19 dering simulation calibration a fundamentally multi-metric optimisation problem 20 that typical calibration techniques cannot handle. Yet calibration is an essential 21 activity in simulation-based science; the baseline calibration forms a control for 22 subsequent experimentation, and hence is fundamental in the interpretation of re-23 sults. Here we develop and showcase a method, built around multi-objective opti-24 misation, for calibrating agent-based simulations against complex target behaviours 25 requiring several metrics (termed *objectives*) to characterise. Multi-objective cal-26 ibration delivers those sets of parameter values representing optimal tradeoffs in 27

simulation performance against each metric, in the form of a Pareto front. We use 28 MOC to calibrate a well-understood immunological simulation against both estab-29 lished a priori and previously unestablished target behaviours. Further, we show 30 that simulation-borne conclusions are broadly, but not entirely, robust to adopting 31 baseline parameter values from different extremes of the Pareto front, highlighting 32 the importance of MOC's identification of numerous calibration solutions. We de-33 vise a method for detecting overfitting in a multi-objective context, not previously 34 possible, used to save computational effort by terminating MOC when no improved 35 solutions will be found. MOC can significantly impact biological simulation, adding 36 rigour to and speeding up an otherwise time-consuming calibration process, and 37 highlighting inappropriate biological capture by simulations that cannot be well 38 calibrated. As such, it produces more accurate simulations that generate more 39 informative biological predictions. 40

41 **1** Introduction

Computational modelling and simulation has emerged as a tool for investigating a wide 42 range of biological systems, spanning immunology [1][2], drug and intervention design 43 [3][4], developmental biology [5], and ecology [6]. Biological simulation is particularly in-44 sightful when used in complement with traditional methods, such as wet-lab in vivo and in 45 vitro work; laboratory work generates experimental data and suggests hypotheses that can 46 be evaluated by way of their integration with simulation, which in turn can suggest further 47 experiments or highlight areas of lacking knowledge [7][8]. Well designed, biologically-48 accurate simulations provide detailed spatio-temporal insight, facilitating observations 49 and assays not possible in the real system; simulation experiments are unhampered by 50 the ethical, practical and financial considerations inherent in biological experimentation. 51 Research programs integrating wet-lab and simulation methods can offer a greater return 52 on animal experimentation by generating additional insight, and hence easing the burden 53 on experimental animals, in line with the '3Rs' principles (Replacement, Reduction and 54 Refinement). 55

The agent-based simulation (ABS) paradigm permits detailed and nuanced simulation of biological systems [9][3]. Simulation components are represented as explicit individual entities, *agents*, with unique states that exist within a spatial environment. Rules ⁵⁹ specifying agent dynamics and the consequences of interaction are provided, and simu-⁶⁰ lation execution allows the system-level consequences of agent-level manipulations to be ⁶¹ observed. ABS incorporates stochastic events, and therein reflects the heterogeneity of ⁶² real world natural systems. There is scope for specifying very detailed interactions using ⁶³ ABS, at the expense of generating large numbers of parameters: 50+ is not uncommon.

Drawing biologically meaningful conclusions from simulation requires that the map-64 ping of the simulation to the biology is known. This can prove problematic for two 65 reasons. First, simulations are abstract representations of their corresponding real world 66 systems. For example, there exist at least 19 varieties of T cell, a vital component of the 67 immune system [10]. However, rather than fully capture all their nuanced differences, a 68 simulation is more likely to represent an abstracted subset thereof. As such, experimen-69 tal measurements on a real world T cell cannot be assumed to translate directly to its 70 simulation counterpart. Second, complex biological systems are the subject of simulation 71 precisely because they are incompletely understood, meaning that the real world data sup-72 porting simulation design decisions and corresponding parameter values may not exist. 73 Calibration is a critical activity in establishing the link between simulation and biology; 74 parameter values that align simulation and real-world dynamics are identified. Further-75 more, an inability to provide a good alignment points to simulation design that does not 76 appropriately capture the biology. Calibration is used to establish a baseline simulation 77 dynamic used as a control in subsequent experimentation, and finding appropriate values 78 is important. Different parameter values will yield different simulation dynamics, and as 79 such influence the conclusions drawn from experiments. 80

A number of approaches to calibration exist, including manual calibration [11], evo-81 lutionary algorithms [12][13], maximum likelihood estimation and various forms of re-82 gression [14]. These techniques identify parameter values by employing a single metric 83 to align simulation dynamics with those of the real world system. However, complex 84 biological system dynamics are not well characterised by single metrics alone. They con-85 stitute many different types of interacting component, and encompass both positive and 86 negative feedbacks. They are highly redundant: a single component can perform many 87 functions and any one function can be performed by several components [15][16]. As such, 88 calibration of a complex system simulation is fundamentally a multi-metric optimisation 89 problem; several metrics of a simulation's alignment with the biology must be simulta-90

neously considered when evaluating putative parameter values. Consider, for example,
cellular motility, which underlies many biological processes arising from cellular interaction. Which targets a given cell interacts with depends on both its speed and directional
persistence; accurately modelling this process requires that metrics of both be considered.

In this paper, we position multi-objective optimisation-based calibration (MOC: multi-95 objective calibration) as an important enabling technology for simulation-based biological 96 investigation. Given its abstractive nature, a simulation undergoing calibration will not 97 perfectly replicate all aspects of the biology. As such, putative simulation parameter value 98 sets will exhibit tradeoffs in their reproduction of aspects of the biology, excelling in some 99 at the expense of others. In this context, a metric quantifying a simulation's capture 100 of a specific aspect of the biology is termed an *objective*. Through the use of Pareto 101 fronts (defined in Section 3), MOC explicitly tracks the collection of simulation parameter 102 sets exhibiting optimal tradeoffs between objectives. It is unknown if adopting baseline 103 parameter values from different regions of the Pareto front will deliver fundamentally 104 different conclusions from simulation-based experiments. The answer to this question is 105 likely problem-specific, and the use of MOC allows this issue to be addressed by exposing 106 a full range of Pareto-equivalent solutions. 107

Here we investigate multi-objective optimisation, specifically the NSGA-II algorithm [17], 108 in calibrating an established immunological simulation: ARTIMMUS [18]. ARTIMMUS 109 simulates Experimental Autoimmune Encephalomyelitis (EAE), a mouse model of multi-110 ple sclerosis [19][20]. It is a complex simulation, encompassing seven distinct cell popula-111 tions that interact across five organs, and constituting 72 parameters. Its successful prior 112 manual calibration renders it an effective test case for evaluating MOC's applicability to 113 simulation calibration. We demonstrate the successful calibration of ARTIMMUS using 114 five objectives (Section 4): a range of solutions to the calibration problem, offering optimal 115 tradeoffs against calibration objectives, are generated. Furthermore, we demonstrate that 116 conclusions drawn from a simulation-based experiment can vary depending on exactly 117 which calibration solution is adopted (Section 5). Hence, different calibration solution 118 parameter values can vary downstream conclusions, highlighting MOC's value in making 119 these multiple solutions explicit. We show that MOC is equally applicable in generating 120 simulation initial condition values: cellular population sizes as simulation launch. We 121 proceed to demonstrate that MOC can identify parameter and initial condition values 122

that deliver previously unknown simulation dynamics, highlighting its potential beyond this well understood test case (Section 6). Lastly, we consider strategies for formulating stopping criteria for MOC, thereby preventing over-fitting and wasted computational expense when apparent improvements in simulation calibration are likely due to stochastic sampling rather than genuinely superior parameter values (Section 7). We begin by introducing ARTIMMUS (Section 2), and the MOC methodology (Section 3).

¹²⁹ 2 A testbed for calibrating biological simulations

ARTIMMUS is an agent-based simulation of an EAE protocol wherein mice induced into 130 autoimmunity undergo a natural recovery from disease, and are thereafter resistant to 131 disease re-induction [18][21][22]. ARTIMMUS was created, in part, to further probe the 132 cellular interactions mediating this recovery [23][24]. It has been used to explore the 133 mechanisms through which splenectomy, the removal of the spleen, a primary immune 134 organ, exacerbates disease severity, and predict the outcome of T cell interaction-blocking 135 drugs [18]. It was conceived through a collaboration of immunologists and computer 136 scientists, and developed through a principled approach focusing on documenting how 137 biological concepts are translated into computer code: the CoSMoS process [32]. It is 138 written in the Java programming language. 139

ARTIMMUS has previously undergone a by-hand, manual calibration [11], and was 140 shown to reflect the dynamics of the real world disease [18]. The process demanded close 141 collaboration between the simulation developer and an immunologist who informed the 142 work, helping bridge biological data and concepts to simulation constructs and output. 143 This manual calibration took two weeks, and entailed an iterative process through which 144 simulation code and parameter value changes that might explain perceived discrepancies 145 between simulation and biological system dynamics were identified and explored in turn. 146 Those best aligning simulation with biological dynamics were adopted before repeating 147 the process. This calibration approach is akin to a non-population, manual, greedy local 148 search wherein the best immediate improvement is always adopted. 149

Despite delivering a well-calibrated result for ARTIMMUS, this calibration search strategy presents several potential pitfalls. It is entirely plausible that the manual search does not find the global optimum parameter set that best aligns simulation dynamics with those of the biological system. As a greedy search strategy, its result is highly dependent on the search's starting position, and complex landscapes where one parameter's influence on simulation dynamics critically depends on the values held by others are particularly challenging. The existence of multiple solutions to the calibration problem can go entirely undetected. Lastly, manual calibration is time consuming, and agent-based simulation's stochastic nature furthers compound these challenges. It is these issues that collectively motivated the present automated MOC approach.

Here we provide a brief summary of EAE and ARTIMMUS to aid understanding of 160 the sections that follow; a comprehensive description may be found in the supplementary 161 materials of [18]. Figure 1A provides an abstract overview of the major cell types in-162 volved in EAE, and their relationships to one another. EAE is induced through injection 163 of neuronal fragments which are internalized by dendritic cells (DCs) which then direct 164 the growth of a T cell population (CD4Th1, abbreviated to Th1) targeting these frag-165 ments. These Th1 cells enter the central nervous system (CNS), where they stimulate 166 CNS-resident macrophages into secreting TNF- α , which in turn damages neurons. The 167 resultant neuronal fragments are internalized by further populations of DCs, which direct 168 further Th1 activities, perpetuating the autoimmune cycle. Recovery from autoimmunity 169 is through the actions of two populations of regulatory T cell, CD4Treg and CD8Treg 170 cells, so named as they regulate the activities of other T cells. The natural life-cycle of a 171 Th1 cell results in its eventual death and internalization by DCs, which derive fragments 172 therefrom and direct the growth of CD4Treg and CD8Treg cells targeting the Th1 cell 173 population. CD4Tregs play an essential role in facilitating the development of CD8Treg 174 cells. CD8Treg cells can directly kill Th1 cells, interrupting their natural life-cycle and 175 preventing the perpetuation of autoimmunity. Th2 cells directly compete with Th1 cells, 176 as both arise from a common progenitor and they each perform downstream activities 177 that promote their own development. The reduced severity of the autoimmune environ-178 ment arising from the action of CD8Treg cells favours the growth of Th2 cells over Th1 179 cells, which do not directly harm neurons and hence do not contribute to this autoimmune 180 process. Figure 1B shows a time-series graph of T cell population sizes in ARTIMMUS. 181

¹⁸² 3 Multi-objective calibration (MOC) methodology

We present here an overview of the multi-objective calibration (MOC) concept, detailing how we employ multi-objective optimisation technology to calibrate simulation parameters and initial conditions. A graphical overview is supplied in Figure 2.

Firstly, we define the desired (*target*) ARTIMMUS dynamics, Figure 2A. In this 186 manuscript targets are expressed as peak cell population sizes, the times at which those 187 peaks occur, or the cell population sizes at a given time. Target dynamics might repre-188 sent known biological results to be reproduced, or hypothetical outcomes of interest. In 189 this study we adopt the dynamics of a previous manual calibration of ARTIMMUS, so 190 as to evaluate MOC on a well-understood problem; thereafter we employ MOC to obtain 191 hypothetical dynamics not known possible *a priori*. We note that many other aspects 192 of simulation performance can constitute target dynamics, depending on the context and 193 simulation being calibrated. The expression of targets as distributions reflects the stochas-194 tic nature of biological systems and agent-based simulations, wherein repeat experiments 195 can yield slightly different results. 196

MOC seeks to identify parameter values that best align simulation with target dynam-197 ics. As such, we define metrics, termed *objectives*, that quantify the alignment between the 198 two. As illustrated in Figure 2B (left), we employ the the non-parametric Kolmogorov-199 Smirnov statistic in our objectives, which quantifies the difference in target and simulation 200 dynamics for a given set of simulation parameter values. Rather than contrasting the me-201 dians of two distributions, as many statistics do, the KS statistic quantifies the biggest 202 distance between two distributions' cumulative distribution functions. As such, its use 203 here facilitates the calibration of a distribution's shape, not simply its median or mean. 204 We consider this a strength of our approach; as may be seen in the sections that follow, 205 MOC is capable of reproducing distributions of behaviour, not simply averages. Each 206 set of simulation parameter values is termed a 'candidate solution', and its corresponding 207 simulation performance is evaluated against each objective individually. By evaluating 208 many candidate solutions we identify regions of parameter space providing close align-209 ment with target dynamics (Figure 2B, right). Importantly, the regions that satisfy each 210 objective differ. In practice, it is computationally intractable to fully explore parameter 211 space as suggested by the heatmaps in this Figure, particularly when many parameters 212 are investigated. Instead, a heuristic (guided) search strategy is employed that samples 213

parameter space, evaluates performance, and decides from where to extract the next can-214 didate solutions based on the results. In this study we employ NSGA-II as our guided 215 search engine [17], but we believe other multi-objective optimisation technologies could 216 be successfully substituted. NSGA-II maintains a population of candidate solutions, and 217 employs (heavily abstracted) principles of genetic recombination, mutation and natural 218 selection to generate and evaluate successive generations of superior candidate solutions. 219 Hence, NSGA-II is an iterative algorithm. We refer the readers to [17] for more detail on 220 NSGA-II. Here we have employed the 'inspyred' python module NSGA-II implementation. 221 We identify those candidate solutions that constitute optimal tradeoffs in performance 222 against each objective, referred to simply as *solutions*, Figure 2C. The set of solutions 223

is termed the *Pareto front*. These solutions are *Pareto-equivalent*: no solution has been 224 found that offers an improvement in one objective without a worsening in another. Pareto-225 equivalent solutions may reside in disparate regions of parameter space, and the ability to 226 recognize this is a key strength of MOC. Though these regions of parameter space may be 227 Pareto-equivalent for the given target simulation behaviour, they could yield very different 228 behaviours when subjected to further downstream experimentation, and as such lead to 220 different simulation-borne conclusions. In this study we investigate this phenomenon for 230 a given experiment in ARTIMMUS. 231

We note that it is possible to derive a great many targets and objectives for complex system simulations. Increasing the number of objectives increases the difficulty of the calibration problem, and the computational resource required to address it; in the field of optimisation this is known as the 'curse of dimensionality'. Hence, employing fewer, uncorrelated objectives is considered good practice: it encourages the identification of good quality solutions whilst minimising the resources required to do so.

²³⁸ 3.1 Selecting candidates from the Pareto front

²³⁹ Upon completion MOC delivers a Pareto front of Pareto-equivalent solutions, representing ²⁴⁰ optimal tradeoffs between the calibration objectives. Deciding which solution adopt as the ²⁴¹ baseline simulation parameter values is an application-specific problem. For the present ²⁴² study we have developed a function, $\Lambda(c)$, which assesses candidate solution c against ²⁴³ the criteria below. We select the candidate with the lowest Λ value when presenting the ²⁴⁴ results of calibration below. Λ is calculated as follows. Let Ω represent the set of calibration objectives, and $KS_o(c), o \in \Omega$ as the corresponding Kolmogorov-Smirnov score for candidate c on objective o. $\overline{KS}(c)$ represents the mean objective score for candidate c. The Λ score is calculated as:

$$\Lambda(c) = \alpha \cdot \overline{KS}(c)^2 + \sum_{o \in \Omega} \left(KS_o(c) - \overline{KS}(c) \right)^2 \tag{1}$$

Low Λ scores are achieved through low mean objective KS scores, and balanced KS scores across all objectives. α specifies the relative importance of these two components. When $\alpha = 1$, both measures contribute equally to Λ . Lower mean KS scores are prioritised with $\alpha > 1$, and vice versa. We employ $\alpha = 1$ throughout. We note that Λ is unit-less, and as such is not explicitly reported here; it is used only to extract one candidate solution from a Pareto front, presented as the chief result of calibration in the results that follow.

²⁵⁴ 4 Successful re-calibration of ARTIMMUS

We demonstrate MOC by re-calibrating ARTIMMUS, taking as target dynamics those of the previous manually-calibrated simulation dynamics [18]. As these dynamics are known to be obtainable, and at least one set of parameter values that produce them are known, we are able to evaluate MOC's performance.

²⁵⁹ With 5 objectives MOC successfully reproduced the manually-calibrated ARTIMMUS ²⁶⁰ dynamics, as demonstrated in Figure 3. The objectives used were:

- the peak Th1 cell population size (Figure 3B)
- the time at which the peak occurred (Figure 3C)
- the Th2 population size at 30 days (Figure 3D)

• the peak population sizes of both CD4Treg and CD8Treg cells (Figures 3E and F).

²⁶⁵ The corresponding target distributions of values are also shown in Figure 3.

Each candidate solution generated by NSGA-II was assessed through 200 replicate simulation executions. The target distributions against which candidates are contrasted are derived from 500 replicates generated with the previous manual-calibration parameter values. The manual-calibration's replicates need be executed once only and stored, they do not change. In contrast, assessment of candidates is computationally costly because

so many are generated; a figure of 200 replicates per candidate was selected to strike a 271 balance between experimental sensitivity and computational cost. A previous analysis of 272 parametric perturbation in ARTIMMUS established that contrasting distributions com-273 prising 200 replicate executions was sufficient to detect 'small' changes in $\frac{2}{3}$ of simulation 274 behaviour metrics, and 'medium' in the remainder [11]. Hence, we consider 200 replicates 275 to offer sufficient sensitivity in differentiating candidate performances. These effect size 276 categories arise from the analysis's use of the Vargha-Delaney A test [25], which provides 277 interpretation guidelines. For reference, the A test is a non-parametric effect magnitude 278 test representing the probability that a randomly selected member of one distribution is 279 larger than a randomly selected member of the other. An A test score of 0.5 indicates 280 the two distributions are indistinguishable (using this test). Values of 1 and 0 indicate 281 no overlap in the two distributions. A single calibration exercise required around 5 days 282 on a dedicated computational cluster able to execute 120 simulations simultaneously; 283 each single simulation replicate takes around 2-10 minutes to execute, depending on the 284 parameter values used. 285

We have successfully applied MOC to both ARTIMMUS parameter values and ini-286 tial conditions, but focus here on the former. Initial condition calibration results are 287 reported in the supplementary materials. Calibration was performed over 8 ARTIMMUS 288 parameters which all pertain to presentation of substances to T cells, particularly Th1 289 and Th2 cells, and their resultant development. The biology captured in these parame-290 ters is outlined in supplementary Figure S1, and we note that a through understanding 291 of this biology is not required to appreciate our results. These parameters were selected 292 for the reasons that ascertaining their values experimentally would be challenging and 293 they all relate to a critical aspect of the biology: the perpetuation of autoimmunity, and 294 (for some) it's amelioration (as Treg cell development is also directed by DCs). Hence, 295 by successfully calibrating parameter values that are highly influential on simulation dy-296 namics we demonstrate MOC's potential. Parameters were given a constrained range of 297 values that the MOC process could assign, being zero to twice their manually-calibrated 298 range, as shown in Table 1. In exploring the space of putative parameter values, NSGA-II 299 maintained a population of 64 candidate solutions which were subject to genetic recombi-300 nation and mutation (see [17]) over 32 generations of natural selection, wherein only the 301 best 64 solutions (i.e. those on or near the Pareto front) were retained in the successive 302

303 generation.

This calibration exercise was repeated three times for both parameters and initial 304 conditions. Figure 3 shows the solution with the lowest Λ score from one such parameter 305 calibration. The remaining two are shown in supplementary Figures S2 and S3. The 306 calibrated simulation dynamics closely resemble the target distributions in all cases. The 307 three parameter calibration exercises generated, respectively, Pareto fronts constituting 308 82, 87 and 112 Pareto-equivalent solutions. The ranges of parameter values represented 309 across the Pareto fronts' solutions in each independent calibration exercise are shown in 310 Figure 4, as are the baseline manually-calibrated values. In all but one case the baseline 311 parameter value sat within the range of non-outlier MOC-derived values, the exception 312 being Th_1_diff80 in exercise 3. Hence, we conclude that MOC is an effective means 313 of calibration: it has repeatedly reproduced ARTIMMUS dynamics that were known 314 possible, and has identified similar solutions, in the form of parameter values, that do so. 315 Next we investigated how the space of ARTIMMUS parameter values relates to the 316 space of successful target dynamic reproductions, i.e., tradeoffs in objective values. We 317 find statistically significant (p < 0.01) differences between calibration exercises' distribu-318 tions of calibrated parameter values for 7 of 8 parameters, Figure 4. This corresponds 319 to 19 of 24 (79%) of pairwise comparisons. Further, 75% (18/24) pairwise comparisons 320 register a KS value ≥ 0.3 . For context, a KS value of 1.0 indicates no overlap between 2 321 distributions. In contrast, this degree of variation is not observed in Pareto fronts' objec-322 tive values, depicted in Figure 5. Here we instead find statistically significant differences 323 in only 27% (4/15) pair-wise calibration comparisons, and only 27% (4/15) of compar-324 isons register KS > 0.3. We find no evidence of objectives that are harder to calibrate 325 than others; the smallest objective values are < 0.05 in all cases, and the median objective 326 values all lie under 0.17. 327

Together, these data suggest a redundancy in the ability for parameter values to deliver particular objective scores. This corresponds to a landscape wherein parameter values mapped to objective values is relatively flat, as a wide range of ARTIMMUS parameter values deliver relatively similar objective scores. The results of using MOC to calibrate ARTIMMUS initial conditions are reported in supplementary Section S1, and supplementary Figures S4, S5 and S6. They are qualitatively identical to our findings in calibrating parameters, and support the conclusions drawn here.

An obvious question is, why does MOC not deliver any perfectly calibrated solutions, 335 wherein all objective scores are 0.0? The best solutions, determined by their minimal Λ 336 values, in each calibration exercise are shown in Table 2. Objective KS values ranged 337 from 0.05 to 0.14 (and 0.03 to 0.12 for initial conditions). We attribute the inability to 338 deliver a perfect calibration to the stochastic nature of ARTIMMUS, wherein 200 replicate 339 executions for a given candidate yields sufficient variation so as to deliver objective KS 340 scores of ≥ 0.05 . There is a risk that improvements in objective KS values that are already 341 so small cannot be confidently attributed to an actual improved simulation calibration, as 342 opposed to stochastic variation between simulation replicates. Section 7, below, explores 343 a method for terminating the MOC process on the premise that further effort will not 344 deliver better quality solutions. 345

These data collectively highlight the challenges in exactly calibrating (i.e. KS=0.0) simulations to several objectives simultaneously. As such, we consider in the next Section the implications on experimental results of adopting baseline simulation values from different extremes of the Pareto front.

5 Scientific significance of imperfect calibration

As demonstrated above, MOC delivers a host of solutions to a given calibration problem, 351 each representing an optimal tradeoff in calibration criteria (see Figure 2). It falls on the 352 simulation developer to decide which to adopt baseline parameter values in subsequent 353 experimentation. There is a risk that whilst calibration solutions lying in different regions 354 of parameter space give rise to Pareto-equivalent solutions, they do not behave in a 355 consistent manner when further experiments are performed. In such a case, a simulation-356 based experiment would lead to different conclusions depending on which calibration 357 result was adopted as the baseline. In this section we investigate the extent to which this 358 phenomenon holds. 359

The manually-calibrated ARTIMMUS simulation was previously used to elucidate the effect of removing a central immune organ, the spleen (a *splenectomy*), in EAE-induced animals [18]. Previous experiments had demonstrated that splenectomy in rats prior to the induction of EAE increased the mortality rate and hampered recovery [26]. Simulating splenectomy in ARTIMMUS revealed the spleen as a primary site for the generation of ³⁶⁵ autoimmunity-combating CD4Treg and CD8Treg cells. The reduced Treg populations
³⁶⁶ resulting from the spleen's removal prior to EAE-induction were unable to completely
³⁶⁷ abrogate the autoimmunity-inducing Th1 populations, allowing for their re-expansion,
³⁶⁸ and thus facilitating increased disease severity and relapses.

Here we explore whether the results of splenectomy in ARTIMMUS differ when base-369 line parameter values are adopted from disparate extremes of the Pareto front. The 370 experimental procedure is highlighted in Figure 6. First, Pareto front solutions represent-371 ing the extreme values, both low and high, of objective KS measures are identified. These 372 solutions represent extremes in the range of simulation dynamics encapsulated within the 373 Pareto front. For each solution 200 simulation replicates are performed for both control 374 and splenectomy groups. Key performance indicators (KPI) are extracted from the resul-375 tant distributions of 200 simulation executions in each group. The performance indicators 376 used are identical to those of the original ARTIMMUS splenectomy experiment [18]: the 377 peak population sizes for each T cell population in the simulation, the times at which 378 these peaks are reached, and the number of Th1 cells remaining at day 40 (giving a total 379 of 9). For each KPI, the distributions of values obtained for control and splenectomy 380 groups are contrasted using the Vargha-Delaney A test [25], as per the original experi-381 ment [18]. This procedure is repeated for each of the three calibration exercises reported 382 in Section 4. The resultant A test scores are shown in Figure 6's tables. Also shown, for 383 context, are the A test scores of the original ARTIMMUS experiment [18]. 384

Broadly speaking, the splenectomy results generated by Pareto-equivalent solutions are 385 consistent with one another, and with the original experiment. There exceptions, however, 386 wherein differences in A test scores reported for solution and the original experiment 387 differed substantially: g23c60 in exercise 1, and g6c35 and g30c58 in exercise 2. These 388 differences occurred for 'Th1 at 40d', 'Th2 peak' and 'Th2 Time' KPIs. Of interest, three 389 of these solutions were obtained from the region of the Pareto front where alignment with 390 target Th2 peak population size was poorest. In the case of g23c60 and g6c35, exercises 1 391 and 2 respectively, the parameter values where sufficient to return Th1 population size at 392 40 days to control group levels, despite the splenectomy (A=0.58 and 0.56; 0.5 indicates 393 no difference). This is significant, as the principle conclusion of the original experiment 394 was that splenectomy reduces Treg population sizes to levels unable to suppress Th1 cell 395 populations and abrogate autoimmunity. The time series T cell population dynamics of 396

³⁹⁷ both these solutions under control and splenectomy are shown in supplementary Figure S9. ³⁹⁸ In both cases the peak Th1 population sizes are smaller than in the original experiment ³⁹⁹ (see Figure 6)), and the Th2 population sizes are substantially larger. Based on this we ⁴⁰⁰ hypothesize that despite reduced Treg population sizes resulting from splenectomy, the ⁴⁰¹ altered balance between Th1 and Th2 populations which compete with one another is ⁴⁰² sufficient to abrogate the Th1 population at day 30 in these solutions.

Supporting the notion that solutions' results are relatively consistent, the direction of change in solutions' KPIs resulting from splenectomy differs from the original experiment in only a minority of cases. Further, this occurs only in KPIs for which the original experiment reports a comparatively small change between splenectomy and control, the largest being in exercise 2 when the original experiment reports a change of A=0.66, which was not interpreted as significant.

We have conducted the same investigation on Pareto-equivalent solutions generated under the three independent initial condition calibration exercises (supplementary Section S1). Detailed analysis is reported in supplementary Section S2 and Figure S10; briefly, divergences between initial condition solution and original experiments were smaller than reported here for parameters. We take this to indicate that the initial parameters investigated were less influential on simulation behaivour than the parameters investigated here.

In summary, the conclusions that would be drawn from adopting baseline parameters 416 values from disparate Pareto-equivalent solutions are mostly, but not completely, consis-417 tent with one another and with the original splenectomy experiment. There were two 418 notable exceptions, and they underscore the importance of considering the range of sim-419 ulation performances that satisfy a calibration exercise. Making these explicit through 420 Pareto fronts is a strength of the MOC approach. It remains important to, where possi-421 ble, further evaluate Pareto-equivalent solutions in the context of domain knowledge and 422 expertise, which might have ruled out the two exceptions noted above, as the Th2 popula-423 tion size is abnormally large compared to the Th1 population. Where this is not possible, 424 where no grounds to discard some Pareto-equivalent solutions exist, we advise that ex-425 periments are performed in replicate adopting a wide range of calibration solutions and 426 that conclusions are drawn after taking stock of the full range of results generated. This 427 is particularly important if quantitative, rather than qualitative, results are sought; our 428

present data show more divergence between calibration solutions and original experiment
in the quantitative case.

⁴³¹ 6 Multi-objective calibration delivers previously un ⁴³² seen disease phenotypes

⁴³³ In Section 4, above, MOC successfully reproduced simulation dynamics known to exist ⁴³⁴ by virtue of a prior manual calibration. To further demonstrate MOC's generality and ⁴³⁵ utility, we now derive simulation dynamics not known to exist *a priori*.

ARTIMMUS's baseline behaviour constitutes a period of autoimmunity followed by 436 recovery, reflecting typical biological disease [21][22]. However, disease susceptibility 437 and severity vary considerably between mouse strains and between mice within a given 438 strain [27][28]. Furthermore, depletion or incapacitation of CD4Treg and CD8Treg cells 439 leads to exacerbated disease symptoms [29][30]. Here we investigate the capacity for 440 ARTIMMUS to reproduce persisting disease symptoms of varying severity. To reflect 441 potential genetic differences between mouse strains, we calibrate over initial conditions 442 specifying cell population sizes, and a parameter controlling the efficiency of Th1 killing 443 by CD8Treg cells; together comprising 9 variables. In this experiment we are implicitly 444 investigating whether variation in these basal population sizes and the efficiency of the 445 CD8Treg-Th1 killing pathways could explain the differences in autoimmune phenotypes 446 observed between mouse strains and individuals therein. 447

Three persisting disease severities are investigated, ranging from mild to severe. These are captured by defining the distribution of Th1 cells remaining at 60 days as a target for calibration, captured as a Guassian distribution. Mild, moderate and severe disease are represented with mean (μ) and standard deviation (σ) values of μ =50 & σ =10, μ =200 & σ =100, and μ =500 & σ =200 respectively. To ensure an aggressive onset of autoimmunity, consistent with animal models, a second calibration target distribution of μ =1000 & σ =200 Th1 cells at 15 days is employed.

Each persisting autoimmunity severity is independently calibrated three times, representatives of which are shown in Figure 7 (the remainder are shown in supplementary Figures S11, S12 and S13). Automated calibration successfully delivers the required median number of cells in most cases, with $KS \leq 0.2$ in 6 of the 9 calibrations. However, the spread of the 'Th1 cells at 60 days' distribution for mild persisting disease is notably less
well calibrated, with all three calibrations delivering KS>0.3.

Together, these data support the general applicability of MOC to problems where a simulation's ability to deliver a desired dynamic is not known *a priori*. These data also suggest that the heterogeneity in disease severities observed in experimental animals could be attributed to differences in basal population sizes and regulatory pathway efficiency.

$_{465}$ 7 When to stop MOC

A key consideration in any optimisation task is the stopping criteria. For MOC, underpinned by the NSGA-II optimisation algorithm, this equates to determining when to stop
calibration.

Overfitting describes the case where the simulation being calibrated starts to capture 469 the noise in the target distributions, rather than the trends those distributions represent. 470 This is a particular issue when target distributions do not contain many samples, as 471 might be the case if they represent biological experiments (Figure 8A). For example, 472 studies involving experimental animals can require their sacrifice to collect data. As such, 473 it is considered unethical (and is practically cumbersome) to collect hundreds of samples, 474 and 5 to 10 are more typical. These smaller sample sizes are unlikely to perfectly capture 475 the underlying distribution that would emerge if thousands of samples were available. 476 Overfitting is said to have occurred when the calibrated simulation better reflects these 477 5-10 samples than their underlying distribution, as illustrated in Figure 8B. 478

A common strategy in single-objective (not MOC, which is multi-objective) problems 479 for determining when to terminate an optimisation process is to segregate the available 480 data into two parts, termed 'training' and 'validation' datasets. The training dataset 481 is used as normal to search for improved solutions, akin to MOC's target data. The 482 validation dataset is used as an independent check for overfitting of solutions to the 483 training data set. Such a case of overfitting is depicted in Figure 8C. Both the training 484 and validation data roughly reflect the underlying distribution, from which they were 485 sampled. The candidate solution more closely resembles the training dataset than either 486 the underlying distribution or the validation dataset, hence, it is overfitted. As illustrated 487 in Figure 8D, in the earlier stages of optimisation successive candidate solutions that 488

better capture the training dataset will also better capture the validation data. It is only when overfitting starts to occur that performance against the validation data worsens whilst performance against training data continues to improve. It is at this point that the optimisation process is best terminated.

⁴⁹³ MOC is, however, a multi-objective optimisation problem, and it is unclear in the lit-⁴⁹⁴ erature how this overfitting detection strategy ought be applied. We propose here a novel ⁴⁹⁵ strategy for detecting overfitting in multi-objective problems based on co-membership of ⁴⁹⁶ solutions to both training and validation dataset Pareto fronts (P_t and P_v), maintained ⁴⁹⁷ throughout the calibration process (Figure 8E). The overfittedness at a given point in the ⁴⁹⁸ optimisation process is reflected in the proportion of P_t members that are not members ⁴⁹⁹ of P_v . The following algorithm performs the calculation:

500 $m \leftarrow 0$ 501 for all $i \in P_t$ do 502 if $i \in P_v$ then 503 $m \leftarrow m + 1$ 504 end if 505 end for 506 return $1 - (m/\text{size}(P_t))$

A proportion of 0 indicates that all training dataset Pareto solutions are also members of the validation Pareto front. At the other extreme, a value of 1 indicates that the training dataset Pareto front has been completely over-fitted, as none of its members are Pareto optimal with respect to the validation dataset. A threshold level of over-fitting at which the optimisation process (i.e., MOC) is to be terminated can be selected by the simulation experimenter.

We investigated different overfitting thresholds for MOC termination in the three 513 ARTIMMUS parameter recalibration exercises reported in Section 4 above. An additional 514 214 simulation replicates using manually-calibrated parameter values were acquired to use 515 as a validation dataset, constituting a 70-30 (500-215) training-validation data split. The 516 validation dataset Pareto front for each iteration of the MOC algorithm (generation) was 517 determined, and the overfittedness calculated. Figure 9A shows how, as MOC progresses, 518 the proportion of overfitted candidate solutions on the training (target) dataset increases 519 for each of the three calibration exercises. Figure 9B shows the point at which MOC 520

calibration would have been terminated should a given overfittedness threshold have been 521 selected. Had we employed a overfittedness termination threshold of 0.5, wherein half 522 of the training dataset Pareto front is overfitted, calibration would have terminated at 523 generation 14, 15 or 23 (for exercises 1, 2 and 3 respectively) instead of 32. Given that 524 each of these calibration exercises required around 7 days to complete on a dedicated 525 computing cluster, this speed-up is substantial. We note that these combined training 526 and validation datasets constitute 714 data points, considerably exceeding what might 527 be obtained from real biological experiments. We anticipate that with fewer data points 528 overfitting will occur sooner in the MOC process. 529

530 8 Discussion

Simulation represents a powerful tool to advance the investigation of biological systems, 531 particularly when used in tandem with traditional approaches. As more complex biolog-532 ical systems become the subject of simulation a challenge in their calibration emerges: 533 complex biological systems cannot be characterised by single metrics alone. There exist 534 technologies capable of identifying parameter values that align simulation dynamics with 535 some desired target, but these operate on single metrics. Even in cases where param-536 eter values can be ascertained experimentally, seemingly avoiding the need for calibra-537 tion, the abstract nature of simulation can complicate their direct adoption. Here we 538 have demonstrated how biological agent-based simulation parameter values can be de-539 rived using multi-objective optimisation, an approach we have termed Multi-Objective 540 Calibration (MOC). Multi-objective optimisation algorithms find solutions to problems 541 simultaneously described by more than one metric. In MOC the desired characteristics 542 of the simulation, which can represent either established biological data to be reproduced 543 or some desired hypothetical simulation outcome, are expressed as distributions. Impor-544 tantly, several such characteristics can be expressed, and MOC identifies those sets of 545 parameter values that deliver optimal tradeoffs against each. 546

We evaluated MOC on a well understood simulation, using it to reproduce a previous manual calibration effort and therein delivering a solution that was known to be possible. The ARTIMMUS simulation was used, which simulates a mouse multiple sclerosis disease model [18]. MOC delivered around 90 unique parameter value combinations,

each of which provided an optimal tradeoff in performance against the 5 target ARTIM-551 MUS characteristics specified. This range of possible calibration solutions was unknown 552 a priori; the previous manual calibration of ARTIMMUS having delivered only one such 553 solution [11]. It would ordinarily fall on the simulation user to select one solution (set 554 of parameter values) to adopt as a baseline for subsequent simulation experimentation. 555 We investigated the significance of selecting solutions representing different extremes of 556 tradeoffs in delivering target simulation characteristics. A previous experiment with AR-557 TIMMUS determined that removing the spleen, an important immune system organ, 558 resulted in exacerbated autoimmune symptoms. The results of re-performing this exper-559 iment with different MOC solutions adopted as baseline parameter values were broadly, 560 but not absolutely, similar. Hence, adopting different calibration solutions can lead to 561 different experimental conclusions. It a strength of MOC that this range of solutions is 562 made explicit. Where possible, we recommend that MOC solutions be evaluated against 563 biological data to discard those that represent biologically unrealistic parameter values or 564 behaviours. Where this is not possible, we advocate performing experiments in replicate 565 using multiple MOC solutions such that the full range of possible results be established 566 before conclusions are drawn. 567

We demonstrated MOC in deriving simulation behaviours that were not known pos-568 sible *a priori*: varying degrees of persisting autoimmunity in ARTIMMUS. MOC can be 569 applied to both parameters and initial conditions, at the same time, as demonstrated in 570 these calibration exercises. We do not consider simulation parameter values and initial 571 conditions as independent; a poor selection of initial condition values coupled with appro-572 priate parameter values can still fail to deliver the desired simulation dynamic. MOC's 573 successful delivery of these previously unknown simulation dynamics presents an inter-574 esting use case for MOC. It could be used to identify which parameters, and hence com-575 ponents and pathways, need be manipulated to resolve a simulated disease state, therein 576 highlighting candidate therapeutic targets. Furthermore, for disease simulations that in-577 corporate potential interventions, MOC can be used to determine optimal intervention 578 strategies that exploit synergies between several treatment options. 579

We surmise that MOC can support model selection and development. Accurately simulating a biological system requires both an appropriate model of the biology, and appropriate parameter values for that model. There typically exist several options for how

to represent a biological concept in simulation, the most suitable of which is often unclear. 583 Models must strike a balance between including sufficient complexity to accurately reflect 584 the biology's dynamics, whilst remaining sufficiently simplistic to offer insight. The un-585 successful calibration of a given model of the biology can lead to two conclusions; first, 586 that the calibration process was simply unsuccessful in finding a solution that does exist, 587 a risk we argue is greatly lessened through MOC; or second, that the model is incapable 588 of replicating the biological dynamics in question. In this latter case, MOC can inform 589 simulation design, where a succession of putative models can be evaluated until calibra-590 tion is successful. The possibility of directly applying MOC to the space of biological 591 abstractions, rather than parameter values, is intriguing, though extremely challenging 592 technically. Here, MOC would search for which cells were represented, and how. This 593 would encompass their interactions with one another, opting to ignore some found to 594 be irrelevant to the biological phenomenon of interest, or *vice versa*. The level of detail 595 through which molecular secretions and expressions where represented could also be de-596 termined; is variable expression level necessary, or does simply 'present' vs 'not' suffice? 597 The challenge herein lies in building an agent-based simulation infrastructure capable 598 of capturing all these possibilities, and allowing the automated optimisation process to 599 manipulate them. The aforementioned point still applies, for each possible model, the 600 space of parameter values must also be investigated, as an accurate reflection of biology 601 requires both an appropriate model and corresponding parameter values. Hence, MOC 602 would be applied in a nested fashion, firstly over the space of biological representations, 603 and therein over the space of parameter values for each model. 604

Although our present investigation has employed an agent-based simulation, MOC 605 is applicable to other simulation paradigms also, such as ordinary differential equations 606 (ODE). Application to non-stochastic simulations, such as ODEs, requires significantly 607 less computational power, as there is no need to obtain simulation replicates in assessing 608 a candidate solution's fitness. We note that, from our experience in building them, not 600 all biological simulations are as computationally costly to execute and calibrate as AR-610 TIMMUS. Each MOC calibration exercise has taken up to a week of time on a dedicated 611 computational facility. In this regard, terminating the MOC process when a threshold 612 level of overfitting is detected is pertinent (see Figure 8). Overfitting was detected in 613 all three of our ARTIMMUS parameter recalibration exercises, and selecting a threshold 614

of 0.5, wherein half of the MOC solutions at a given point no longer represent optimal
performance tradeoffs in an independent test, could as much as halve the computational
effort required.

The ability to detect overfitting in a multi-objective context is a novel contribution 618 of this work. Though a common strategy for stopping a single-objective optimisation 619 process, it was previously unclear how to deploy this strategy in a multi-objective context 620 [31]. There is another condition under which we feel it pertinent to terminate the MOC 621 process. The goal of MOC is to find parameter values yielding simulation dynamics that 622 closely resemble some target. As this alignment increases, and differences in solutions' 623 simulation performances reduce, it is possible that seemingly better alignments in fact 624 represent sampling artefacts arising from the stochastic simulation, rather than genuinely 625 superior parameter values. We note that detecting this in a statistically robust manner 626 is challenging, and as such we highlight it as potential further work. 627

This work fits within the context of a wider framework for supporting complex system 628 simulation, the CoSMoS framework [32]. CoSMoS advocates explicitly recording, typically 629 through graphical modelling [33], how real world concepts are translated into computer 630 code, and the implicit assumptions therein. In this context, MOC can help in relating 631 simulation results to biological data. The case where a distribution of results emerges 632 from a given biological experiment, even to the point where replicates or individuals 633 within an experiment exhibit completely different outcomes, can be handled in MOC 634 by defining bi-modal (or multi-modal) target distributions. A scenario wherein MOC 635 unexpectedly delivers several distinct and unconnected simulation phenotypes, rather 636 than a continuum of points on the Pareto front, is interesting. This can either suggest 637 the existence of additional phenotypes to look for in the biology, or if this can be ruled 638 out, suggests instead that the model being calibrated fails to accurately capture the 639 biology. This later case is an example of how MOC could drive simulation design and 640 development, as covered above. Related work on supporting the link of simulation to 641 biology proposes the construction of an argument wherein a claim such as 'this simulation 642 is an adequate representation of the biology' is supported by explicitly cited evidence [34]. 643 In this context, application of MOC can raise confidence that appropriate parameter and 644 initial condition values have been identified. The range of possible values can be contrasted 645 against biological literature and data, excluding those deemed implausible. Subsequent 646

⁶⁴⁷ simulation experiments can be performed in replicate with those that remain, therein
⁶⁴⁸ highlighting the full range of results that are plausible in absence of better reason to rule
⁶⁴⁹ out particular parameter values. We argue that drawing conclusions from this nature of
⁶⁵⁰ simulation experimentation, and making explicit the full range of parameter values that
⁶⁵¹ satisfy the calibration problem, leads to more robust conclusions.

In summary, our novel application of multi-objective optimisation in MOC presents the multi-objective optimisation community with a new field of application, and one we feel has considerable scope for growth. Importantly, it provides fundamental support for a critical aspect of simulation-based biological experimentation: identifying parameter values and initial conditions that align simulations with a complex target behaviour.

657 Competing interests

⁶⁵⁸ We have no competing interests.

Authors' contributions

MR conceived and designed the study, carried out all data generation and analysis reported herein, and drafted the manuscript. KA conceived the study and drafted the manuscript. LR conceived the study. JT conceived the study and drafted the manuscript.

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Figure and table captions

Figure 1. The ARTIMMUS simulation, used as a testcase for evaluating MOC. 759 **A**, the the major cell types represented in ARTIMMUS, and their key influences on one 760 another. Red and green arrows respectively indicate activities that perpetuate autoimmu-761 nity or mediate recovery. Figure adapted from [11]. **B**, the baseline dynamic of ARTIM-762 MUS, depicting four T cell population sizes over time. The simulation behaviour depicted 763 here forms a calibration target for MOC in Section 4. Lines correspond to like-coloured 764 cells in Figure A; these colours are maintained throughout the manuscript. Error bars 765 capture 90% of the data derived from 500 simulation executions, timeseries lines indicate 766 median population sizes at each time point. 767

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Figure 2. Overview of the Multi-Objective Calibration (MOC) concept. A, 770 The desired (target) simulation dynamics are defined as distributions (only 2 shown): the 771 desired distributions of peak cell number and the times at which these occur. Distribu-772 tions are depicted as histograms, or the corresponding cumulative distribution functions 773 describing the proportion of samples in the distribution (y axis) that hold a given value 774 or less (x axis). **B**, the capacity for putative simulation parameter (only 2 shown) values, 775 termed candidate solutions, to reproduce target dynamics is evaluated. The Kolmogorov-776 Smirnov (KS) statistic quantifies the difference between target and a given candidate 777 solution's simulation performance (left); this metric is termed an *objective*. By sampling 778 and evaluating regions of parameter space we identify those that provide good alignment 779 with a given objective, illustrated through greyscale heatmaps (right). No single region 780 of parameter space maximizes performance against all objectives (only 2 shown), there 781

exist inherent tradeoffs. A heuristic (guided) search strategy, NSGA-II, is employed to strategically sample parameter space. **C**, *solutions* representing optimal tradeoffs in performance against each objective are identified, collectively termed the *Pareto front* (left). These solutions are *Pareto-equivalent* (pink): no solution has been found that represents an improvement in one objective without a worsening in another. Sub-optimal candidate solutions are discarded (blue). Pareto-equivalent solutions may reside in disparate regions of parameter space(right).

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Figure 3. Multi-objective calibration (MOC) successfully re-calibrates AR-791 TIMMUS parameters against 5 objectives. The best solution's, that with lowest 792 A score, target simulation dynamics are shown. The solution dataset comprises 200 sim-793 ulation replicates, the target comprises 500. A, T cell population sizes over time, for 794 both target (dotted line) and solution (solid line). The median values from each dataset 795 at the given point in the time series are plotted. B-F, cumulative distribution functions 796 showing alignment of solution and target distributions of values for each objective, with 797 titles giving KS values. These graphs show the distribution of calibration target values 798 obtained in each dataset: the y-axis indicates the proportion of items in the distribution 799 holding a value less than or equal to the corresponding x-axis value. Objectives are: B, 800 peak CD4Th1 population size cell; C, time at which this peak occurs; D, CD4Th2 pop-801 ulation size at 30 days; E, peak CD4Treg population size; F, peak CD8Treg population 802 size. These data represent the first of three independent recalibration experiments. 803

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Figure 4. Automated re-calibration of ARTIMMUS parameters delivers so-806 lutions approximating the original manually-calibrated parameter values. Box 807 plots are shown for each of three independent calibration exercises. The horizontal green 808 line represents the manually-calibrated parameter values. Calibration was performed over 800 5 objectives: the peak population sizes of Th1, CD4Treg, CD8Treg cells, the time at which 810 the Th1 population peaks, and the number of Th2 cells at 30 days. Parameters subject 811 to calibration are listed in Table 1, see Figure S1 for an explanation of their operation in 812 ARTIMMUS. Values shown above each plot are the Kolmogorov-Smirnov scores between 813

distributions, shown to one significant figure; the associated p-values are: *, p<0.01 and **, p<0.001. Outliers in boxplots are defined as lying beyond the first or third quartiles by 1.5 times the interquartile range.

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Figure 5. The range of objective values that constitute the Pareto front de-819 rived through MOC re-calibration of ARTIMMUS. Box plots are shown for each of 820 three independent calibration exercises. These objective values correspond to the Pareto 821 front and associated ARTIMMUS parameter values of Figure 4. Calibration was per-822 formed against five objectives: \mathbf{A} , the peak population size of Th1 cells; \mathbf{B} , the time at 823 which this occurred; C, the number of Th2 cells at 30 days; D, the peak population size 824 of CD4Treg cells; E, the peak population size of CD8Treg cells. Statistical and boxplot 825 formatting are as in Figure 4. 826

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Figure 6. Do different regions of MOC's Pareto front of solutions give rise 820 to different results in subsequent experimentation? Top, an overview of the ex-830 perimental procedure. 1, Pareto front members representing objective value extremes 831 are identified (only two objectives shown in example). 2, The simulation parameters 832 represented by such members are adopted in performing a control and splenectomy ex-833 periment, with 200 replicate simulations in each group. 3, Key performance indicators are 834 extracted from the resultant distributions of simulation dynamics, indicated here is the 835 peak CD4Treg population size within each individual simulation. 4, Performance indica-836 tors are statistically contrasted for splenectomy and control experiments. These statistics 837 are examined across different Pareto front members, thereby gauging the extent to which 838 experimental results critically depend on which Pareto-equivalent parameter values are 839 adopted in simulation. **Tables**, columns represent extreme Pareto front solutions, defined 840 as having either the highest or lowest KS value for each of the five objectives used in cal-841 ibration (see Section 4). The objective KS value scores are shown in parentheses. Only 842 the first occurrence of each solution is shown, with subsequent entries indicated by '-'. 843 Rows indicate the difference between control and splenectomy simulations based on each 844 solution according to key indicators of simulation behaviour, as measured by the Vargha-845

Delaney A test [25]. The original A test scores for the manually-calibrated simulation 846 are shown ('orig'), as is the biggest difference in A test score observed between manually-847 and automatically-calibrated simulations ('diff'). Values highlighted in red represent four 848 differences in candidate and original A test scores that are notably larger than differences 849 observed elsewhere. 'D.c.' indicates 'direction change', where there exists at least one 850 candidate with for which the A test score lay on the other side of 0.5 from the original, 851 indicating that the distribution of values under splenectomy increased in the original ex-852 periment but decreased for the candidate (or vice versa). 853

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Figure 7. Employing MOC to discover parameter and initial condition val-856 ues delivering simulation dynamics not known to exist a priori: persisting 857 autoimmune states of varying severity. Three severities are explored, represented 858 as columns. They are, a mean of 50, 200 or 500 Th1 cells at 60 days (with standard 859 deviations of 10, 100 and 200 respectively). A second objective is employed in all cases, 860 1000 Th1 cells at 15 days, which drives the establishment of autoimmunity. Each severity 861 is calibrated in three independent experiments, and shown here are the solutions exhibit-862 ing lowest Λ values from a representative calibration of each experiment. The first row 863 of graphs depicts the median T cell time-series. The second row shows the candidate's 864 performance against an objective of 1000 Th1 at 60 days (standard deviation = 200). The 865 last row depicts the second objective, the (respective) number of T cells at 60 days. 866

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Figure 8. Terminating MOC when overfitting occurs. Overfitting describes the 869 case when solutions generated by an optimisation process, e.g. MOC, better resemble 870 the target data than the underlying distribution from which it was drawn. A, in many 871 contexts, such as animal experiments, only limited samples of a phenomenon can be ob-872 tained. The samples will broadly, but not exactly, reflect the underlying distribution. **B**, 873 an overfitted candidate solution more closely resembles the target data than the underly-874 ing distribution from which the target data was drawn. Detecting this is difficult because 875 the true underlying distribution cannot be absolutely known. C, a common strategy in 876 single-objective optimisation problems is to divide the available data into two, a training 877

dataset and a validation dataset. The training dataset is used as the target in obtaining 878 successively better quality solutions. The validation dataset is used as an independent 879 check. Overfitting is detected when solutions more closely resemble the training dataset 880 than validation dataset. This is illustrated in **D**, where early solutions generally offer 881 improved performance against both datasets. It is only in later stages that solutions so 882 closely reflect the target dataset that they diverge from the validation dataset. This is 883 when the process should be stopped. E. Overfitting can be detected in multi-objective 884 optimisation, such as MOC, by maintaining Pareto fronts of optimal solutions against 885 both training and validation data independently. The degree of overfitting is reflected 886 in the proportion of training data Pareto front solutions that are not members of the 887 validation data Pareto front. 888

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Figure 9. Empirical results for detecting overfitting in MOC, and when to 891 terminate the process accordingly. We generated a validation dataset using AR-892 TIMMUS's previous manually-calibrated parameter values, and retrospectively analysed 893 how overfitted MOC solutions would have been on the three MOC calibration exercises 894 reported in Section 4. A, The overfittedness, defined as the proportion of MOC Pareto 895 front solutions that are not also members of a similar Pareto front maintained for the 896 validation data, at each MOC generation. **B**, the generation at which MOC would have 897 been terminated for a given overfittedness threshold value. 898

899 900

904

905

Table 2. The best solution, being that with the lowest Λ value, arising from each of three independent calibration exercises. Shown are each of the five objective KS values. We independently investigated the calibration of both ARTIMMUS parameters (top) and initial conditions (bottom). High quality calibrations, as indicated by low KS values, were

<sup>Table 1. The ARTIMMUS parameters (top) and initial conditions (bottom) subject to
calibration, their baseline (manually-calibrated) values, and the lower and upper bounds
of values they may be assigned during MOC.</sup>

⁹¹⁰ obtained in all cases.

⁹¹¹ Figures and tables



Figure 1:



A. Define target simulation dynamics

Time B. Evaluate putative simulation parameters' reproduction of target dynamic



'Candidate solutions' of putative parameter values identified through heuristic (guided) search: NSGA-II.

Target 1: distribution of peak cell number, across



C. Identify Pareto-optimal solutions (the Pareto front)



Location of Pareto-equivalent solutions in parameter space



Figure 2:



Calibrate ARTIMMUS parameters, exercise 1

Figure 3:



Figure 4:



Figure 5:



Calibration Exercise 1, Vargha-Delaney A Test Scores													
	Th1Peak		Th1Time		Th2at30d		CD4TregPeak		CD8TregPeak				
	g31c32	g11c33	g29c37	g10c13	g31c6	g23c60	g30c0	g23c22	g10c13	g23c22	orig	diff	d.c.
Response	(KS=0.04)	(0.81)	(0.03)	(0.60)	(0.03)	(0.73)	(0.04)	(0.65)	(0.04)	(0.51)			
Th1 Peak	0.70	0.73	0.65	0.67	0.67	0.66	0.71	0.67	-1	-	0.62	0.11	
Th1 Time	0.53	0.51	0.55	0.42	0.53	0.52	0.52	0.52	-	-	0.47	0.08	Y
Th2 Peak	0.60	0.74	0.66	0.66	0.67	0.67	0.61	0.62		-	0.66	0.08	
Th2 Time	0.47	0.53	0.68	0.46	0.51	0.56	0.53	0.47	-	-	0.58	0.11	Y
CD4Treg Peak	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	-		0.00	0	
CD4Treg Time	0.23	0.24	0.26	0.26	0.20	0.28	0.20	0.24	-	-	0.21	0.07	
CD8Treg Peak	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	-	-	0.00	0	
CD8Treg Time	0.24	0.24	0.29	0.27	0.23	0.25	0.25	0.26	-	-	0.23	0.06	
Th1 at 40d	0.98	0.96	0.92	0.89	0.96	0.58	0.96	0.94	-	-	0.95	0.37	

Calibration Exercise 2, Vargha-Delaney A Test Scores													
	g30c25	g6c35	g30c34	g9c63	g15c14	g30c58	g17c61	g14c54	g9c56	g14c54	orig	diff	d.c.
Response	(KS=0.03)	(0.94)	(0.03)	(0.33)	(0.04)	(0.98)	(0.04)	(0.77)	(0.04)	(0.67)			
Th1 Peak	0.63	0.69	0.66	0.66	0.71	0.65	0.65	0.66	0.68	<u>-</u>	0.62	0.09	-
Th1 Time	0.53	0.49	0.55	0.47	0.50	0.53	0.54	0.51	0.48	-	0.47	0.08	Y
Th2 Peak	0.66	0.67	0.65	0.70	0.68	0.39	0.65	0.61	0.70	-	0.66	0.27	Y
Th2 Time	0.56	0.51	0.63	0.63	0.50	0.31	0.54	0.48	0.54	-	0.58	0.27	Y
CD4Treg Peak	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	-	0.00	0.01	
CD4Treg Time	0.27	0.30	0.23	0.19	0.20	0.21	0.17	0.22	0.29	-	0.21	0.09	
CD8Treg Peak	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00	0.00	-	0.01	0.01	
CD8Treg Time	0.27	0.26	0.23	0.17	0.21	0.20	0.22	0.23	0.28	-	0.23	0.06	
Th1 at 40d	0.96	0.56	0.94	0.91	0.94	1.00	0.97	0.97	0.82	-	0.95	0.39	

Calibration Exercise 3, Vargha-Delaney A Test Scores													
	g5c39	g23c10	g28c47	g13c21	g11c6	g5c39	g24c62	g23c10	g23c40	g23c10	orig	diff	d.c.
Response	(KS=0.03)	(0.99)	(0.04)	(0.48)	(0.04)	(0.82)	(0.03)	(0.98)	(0.04)	(0.94)			
Th1 Peak	0.65	0.65	0.63	0.70	0.68	-	0.65	-	0.70	-	0.62	0.08	
Th1 Time	0.43	0.49	0.49	0.42	0.47	-	0.47	-	0.52	-	0.47	0.05	Y
Th2 Peak	0.67	0.64	0.70	0.68	0.67	-	0.67	-	0.67	-	0.66	0.04	
Th2 Time	0.59	0.51	0.52	0.50	0.47	-	0.62	-	0.57	-	0.58	0.11	Y
CD4Treg Peak	0.07	0.02	0.00	0.00	0.00	-	0.00	-	0.00	-	0.00	0.07	
CD4Treg Time	0.30	0.35	0.21	0.24	0.26	-	0.20	-	0.19	-	0.21	0.14	
CD8Treg Peak	0.04	0.01	0.00	0.00	0.00	-	0.00	-	0.00	-	0.00	0.04	
CD8Treg Time	0.27	0.40	0.17	0.26	0.27	Ξ.	0.20	-	0.21	-	0.23	0.17	
Th1 at 40d	0.93	0.83	0.97	0.90	0.92	-	0.97		0.95	-	0.95	0.12	

Figure 6:



Figure 7:



Figure 8:



Figure 9:

Parameters calibrated									
Parameter	Baseline value	Lower bound	Upper bound						
$APC_{-}immatureDuration$	48	0	96						
$APC_matureDuration$	110	0	220						
$APC_phagocytosisToPeptide$	0.02	0	0.04						
CNSM_MBPExpressionProbability	0.2	0	0.4						
$DCT1_cytokineSecretionRate$	10	0	20						
$DC_{-}T2CytokineRatio$	0.17	0	0.34						
$Th1_diff00$	0.05	0	0.1						
$Th1_diff80$	0.85	0	1.0						
Initial conditions calibrated									
Initial condition	Baseline value	Lower bound	Upper bound						
numTh	40	0	80						
numCD4Treg	30	0	60						
numCD8Treg	30	0	60						
numCNS	500	0	1000						
num CNSM a crophage	75	0	150						
numDC	10	0	20						
numDCCNS	40	0	80						
numDCSnleen	100	0	200						

Table 1:

Calibration on parameters										
Calibration	Objective KS value									
exercise	Th1Peak	Th1Time	Th2at30d	CD4TregPeak	CD8TregPeak					
1	0.06	0.10	0.08	0.06	0.07					
2	0.08	0.06	0.06	0.05	0.07					
3	0.05	0.08	0.14	0.08	0.05					
Calibration on initial conditions										
Calibration	Objective KS value									
exercise	Th1Peak	Th1Time	Th2at30d	CD4TregPeak	CD8TregPeak					
1	0.06	0.08	0.04	0.03	0.06					
2	0.04	0.08	0.10	0.11	0.12					
3	0.06	0.06	0.06	0.07	0.05					

Table 2: