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Different reprogramming propensities in plants and mammals: Are small variations in the core network wirings responsible?

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Abstract

Although the plant and animal kingdoms were separated more than 1.6 billion years ago, multicellular development is for both guided by similar transcriptional, epigenetic and post-transcriptional machinery. One may ask to what extent there are similarities and differences in the gene regulation circuits and their dynamics when it comes to important processes like stem cell regulation. The key players in mouse embryonic stem cells governing pluripotency versus differentiation are Oct4, Sox2 and Nanog. Correspondingly, the WUSCHEL and CLAVATA3 genes represent a core in the Shoot Apical Meristem regulation for plants. In addition, both systems have designated genes that turn on differentiation. There is very little molecular homology between mammals and plants for these core regulators. Here, we focus on functional homologies by performing a comparison between the circuitry connecting these players in plants and animals and find striking similarities, suggesting that comparable regulatory logics have been evolved for stem cell regulation in both kingdoms. From in silico simulations we find similar differentiation dynamics. Further when in the differentiated state, the cells are capable of regaining the stem cell state. We find that the propensity for this is higher for plants as compared to mammalians. Our investigation suggests that, despite similarity in core regulatory networks, the dynamics of these can contribute to plant cells being more plastic than mammalian cells, i.e. capable to reorganize from single differentiated cells to whole plants—reprogramming. The presence of an incoherent feed-forward loop in the mammalian core circuitry could be the origin of the different reprogramming behaviour.

Introduction

The differences in reprogramming competence between plants and animals might not be surprising, as the survival of plants requires more plasticity regarding shape and form given their fixed location. However, the underlying mechanisms for these differences are still obscure.
Understanding the origin of these differences would be very elucidating when comparing the different kingdoms and could be of paramount value for further development of efficient reprogramming recipes in mammalian systems with impact on tissue regeneration in health care. The origin of these differences between the kingdoms could be of transcriptional, epigenetic or signalling nature or combinations thereof. In here we focus on transcriptional regulation as the available epigenetic and signalling data do not provide a clear answer to our question.

To this end we will consider the in silico dynamical properties of the core transcriptional architectures for Oct4, Sox2, Nanog [1–3] inside embryonic stem cells (ESC) and WUSCHEL (WUS), CLAVATA3 (CLV3) for shoot apical meristem (SAM) respectively [4–6]. We put forward computational models for ESC and SAM exhibiting bistability. The two stable states are linked to the differentiated and pluripotent cell state respectively. We envisage reprogramming and differentiation as dynamical transitions between the two stable states, and analyse the differences in the transition dynamics between the mammalian and plant networks.

Prior to presenting the dynamical models and their results, we briefly review what is known from epigenetics and cell signalling in the context of plant and mammalian stem cell differentiation and reprogramming.

Chromatin modifications have been identified as important for differentiation and reprogramming. Chromatin remodelling is key to reprogramming in mammalian stem cell systems [7, 8]. In the core Oct4/Sox2/Nanog architecture for embryonic stem cells (ESC) Oct4 opens up the Nanog region by turning on the Jmjd1a and Jmjd2c demethylases [9]. The H3K9me3 mark is responsible for making the chromatin of most pluripotency genes inaccessible for binding. The demethylases activated by Oct4 lead to global H3K9me3 depletion improving cell reprogramming [10, 11]. Nanog is important for the ESC state as it facilitates transition of partially reprogrammed cells towards ground state [12]. Moreover, Nanog enhances the expression of Oct4 by recruiting Tet1, which modulates DNA methylation levels at CpG-rich promoters, promotes transcription of pluripotency factors and participates in the repression of Polycomb-targeted developmental regulators [13, 14].

In the Shoot Apical Meristem (SAM), stem cells are maintained throughout the life of the plant. Cells are stuck together via cell walls and stem cells situated at the very apex are pushed, via growth, to the periphery of the apex where differentiation starts. At the core of the regulation of stem cell maintenance is the homeodomain transcription factor WUSCHEL (WUS) sufficient and necessary for stem cell activity [15]. WUS is expressed in the central part of the meristem, can move between cells [16, 17], and has been shown to activate stem cell genes and repress (potentially with co-factors) differentiation genes [18]. WUS activates CLAVATA3 (CLV3) in the stem cells, which is part of negatively regulating WUS. Together, this forms a negative feedback regulation assumed to be at the core of the stem cell maintenance regulation. In flower primordia there is a transient stem cell activity regulated by the same CLV3/WUS network as in the SAM before differentiation into specialized flower organs [4]. The inactivation of WUS expression here results from an interplay between transcriptional and epigenetic regulation [19, 20]. While factors with chromatin remodelling effects within the SAM have so far been identified [21, 22], no specific mechanistic relationships with the core network as for the mammals have so far been mapped out. This asymmetry in detailed knowledge of epigenetic regulation between the two kingdoms could be due to the surge in focus when it comes to reprogramming with regenerative medicine as a goal.

In the context of stem cell reprogramming, also external signals have been proposed to be important [23–26]. Plant cells react to relative levels of the plant hormones auxin and cytokinin for regeneration of shoot or root tissues from cells extracted from different organs [23]. The plant protocol does not require manipulations in terms of external over-expression of
certain genes in the differentiated cell in contrast to the mammalian case [24]. In the SAM, cytokinin has been suggested to directly impact the CLV3/WUS system to activate the central stem cell core [27], while high auxin levels are important for the initiation of new organs at the periphery [28]. Also microRNAs have been proposed as a main intercellular signalling molecule for several developmental processes in plants, including the regulation of stem cell maintenance [25]. In short, plant stem cell regulation has been evolved to heavily make use of intercellular communication within its niche. So far, transcription factor transfer between cells has not been identified in mammalian cells, though it has been shown that Oct4 forms a complex with E-cadherin and β-catenin at the membrane of the mouse ESC playing a central role in pluripotency [26]. In conclusion, plants’ needs for responding to the environment, partly connected to signalling molecules and lack of cell migration, can be connected to a more abundant use of cell-to-cell communication compared to animals. Whether this is due to a more complex extracellular matrix in the latter or this is enough to explain differences in plasticity remains to be understood.

Results

Direct comparison between the core regulatory networks in plants and mammals identifies similarities

**The shoot apical meristem model.** We define a model using the main components of the SAM dynamics (Fig 1A). For the stem cell activity these are three CLAVATA genes (CLV1, 2 and 3) [6, 29, 30], and WUSCHEL [31]. CLV3 is localized in the central zone [27] and is recognized as a marker of stem cell identity whereas WUS is expressed below in the organizing centre [13]. The three CLV genes have been suggested to function in the same pathway to control meristem development [5, 29, 30] by repressing WUS [32, 33]. WUS, on the other hand, induces the expression of CLV3 [33] and thereby stem cell identity.

GSAM represents the genes expressed in the peripheral zone in the differentiated cells. In this study, the interactions for GSAM are based on KAN1, since perturbation data are available for this gene [18]. However several genes behave in a similar manner, some of which were analysed for WUS changes in [18]. The network topology built based on these experimental observations is shown in Fig 1A. It should be noted that in here we exclude all transports between expression domains in the SAM model and focus on the intracellular expression states.

**The embryonic stem cell model.** In the embryonic stem cell the core of the gene regulatory network consists of the genes OCT4, SOX2 and NANOG [1–3]. These three transcription factors maintain the pluripotent state. The regulation of the three main genes by the transcription factors OCT4 and SOX2 is attained through the formation of the two into a heterodimer—an OCT4/SOX2 complex [34]. OCT4/SOX2 induces its own transcription [34] whereas NANOG, according to recent findings, represses its own expression [35, 36]. In [9] an epigenetic effect is suggested in which OCT4 activates a component that opens up, among other genes, NANOG. This means that the regulatory region of NANOG is inaccessible in the absence of OCT4. Other than the three genes mentioned above the OCT4/SOX2 complex also induces FGF4, which promotes differentiation through NANOG repression [37]. This model is presented in [38] and it also includes differentiation gene G. Possible candidates for gene G are Sox17 and Gata6.

In this study, we propose a simplified ESC network compared to the one in [38] where we remove FGF4 and include instead direct repression of NANOG by OCT4/SOX2 complex (Fig 1A).

Assuming that OCT4/SOX2 translates into CLV3, NANOG into WUS and GESC into GSAM a comparison between the models reveals that three interactions, marked in black in Fig 1A,
Fig 1. Comparison between network topology and differentiation dynamics of the two single cell minimal models of SAM [18] and ESC [38]. A) Gene regulatory networks where black interactions are common to both models, purple interactions are specific for ESC and the orange interactions indirectly exist in the plant stem cell (SAM) dynamics. For model and parameters descriptions see Methods section. B) Examples of time series results of differentiation simulation with SAM and ESC model.

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are common to both models. All self-regulatory interactions and induction of NANOG by OCT4/SOX2 are specific to the ESC model (purple in Fig 1A) and the interactions between $G_{ESC}$ and OCT4/SOX2 are indirect reactions in the SAM model (orange in Fig 1A). The repression of OCT4/SOX2 by $G_{ESC}$ exists in the SAM network through $G_{SAM}$ represses WUS, which activates CLV3, while CLV3 activates $G_{SAM}$ by repressing WUS, which is a $G_{SAM}$ repressor. Similarly, the direct activation of the CLV3 from WUS in the plant network is included in the ESC system by NANOG repressing the repressor of OCT4-SOX2. In summary, we find striking similarities between SAM and ESC model with regard to majority of the regulatory interactions if indirect interactions are accounted for although the ESC system has a few additional identified regulatory interactions.

**Dynamical properties of transcriptional stem cell regulation suggest differences between plants and mammals in transition propensities from differentiated to stem cell states**

The similarities in molecular mechanisms identified and the structure of the transcriptional regulatory modules cannot explain the differences in reprogramming competence between plants and mammals. Hence it is tempting to pursue dynamical modelling in order to identify such differences due to the complexity resulting from the differences of interactions (Fig 1A).

Using simplified stochastic computational models for the mammalian [38] and plant stem cell [18] dynamics on the single cell level reveal great similarity in the dynamical behaviour of the systems (Methods, Fig 1B). The single cell approach is a simplification, as intercellular signalling within the stem cell niches are replaced by externally applied signals, but still captures the main dynamics and attractors of the systems [18, 38]. In both the animal and plant cases, the systems have natural differentiation dynamics where they spontaneously jump between the stem cell state and the differentiated state, while they less frequently (typically never in our long simulations) spontaneously jump from the differentiated state to the stem cell state (Fig 1B). This holds true for a range of parameter values selected for additional independent perturbation dynamics (Methods, [18, 38]). The similarity may reflect the core mutual inhibition between stem cell factors and differentiation factors described above that has been identified in both animals and plants [18, 38].

Given this similarity it is of interest to also investigate how the circuits behave when simulating reprogramming experiments, within the parameter ranges of the models. The ESC and SAM models give rise to very different reprogramming behaviour, Fig 2. The SAM model efficiency has a sharp transition from zero to one, where one means that all simulations led to successful reprogramming. The ESC model, on the other hand, has a reprogramming efficiency that peaks for some reprogramming forces and then decreases. This corresponds well to the experimental and computational result that reprogramming works best for a limited range of over-expression [38, 39, 40]. In plants, ectopic WUS, either directly or via removing the function of CLV3 has been used to create ectopic stem cells in vivo [16, 41, 42]. In the ESC system over-expressing Oct4 would represent reprogramming experiments in the simplified model [43, 44, 45]. Strikingly, the plant system reaches its maximum in de-differentiation earlier as compared to the mammalian system (Fig 2). More importantly, the drop in the latter for high over-expression is consistent with experimental results [38, 39] and is a consequence of the attractor structure of the non-linear dynamics for this model (Fig 2B). While this might indicate a failure of de-differentiation at high levels or a push of the system into another differentiated state, it clearly predicts that a more specific treatment is needed for de-differentiation in the mammalian system [18, 38].
Fig 2. Reprogramming efficiency by over-expressing Oct4 and WUS respectively. Over-expression was implemented by adding a constant production (reprogramming force) to the equation for OCT4/’SOX2 mRNA rate of change in the ESC model and correspondingly for WUS in the SAM model. If the addition of this constant resulted in a switch of the system to a pluripotent state we considered the reprogramming to be successful. The results are based upon monitoring the reprogramming success in 100 independent stochastic Gillespie runs. A) Reprogramming efficiency for different levels of WUS overexpression in the plant SAM model. (B) Reprogramming efficiency for the

C

ESC for different $p_3$ values

$\star p_3 = 1.5$

$\star p_3 = 0.9$

$\star p_3 = 0.2$
In summary, our simplified regulatory network models indicate that although they have similar interactions and differentiation dynamics, the models capture the increased difficulty of reprogramming in the ESC cells compared to plant stem cells.

Adjusting the interaction strengths in the ESC network can increase the propensity for reprogramming

Furthermore, we attempted to elucidate which couplings in the ESC network that might be responsible for the difference in reprogramming efficiency between the SAM and ESC systems. To this end we performed simulations where we modified one parameter at a time while keeping the others unchanged and assessed the impact on reprogramming efficiency.

The parameters that influenced the most the reprogramming outcomes are $p_1$ and $p_3$ corresponding to the incoherent feed-forward motif for Oct4 regulating Nanog (Table C in S1 File). If $p_1$ is reduced, corresponding to the activating interaction, Nanog cannot be induced by Oct4 and reprogramming is lost. The parameter $p_3$ is representing the Nanog repression by Oct4. Fig 2C shows reprogramming efficiency for various levels of the reprogramming force when Oct4 represses Nanog with three different strength values. The results show that reducing Nanog repression by Oct4 leads to an ESC reprogramming efficiency similar to that of SAM (Fig 2C green data). Of note is that Oct4 repression of Nanog can be indirect through e.g. Fgf4. However, in this study we considered a simplified network topology and assume that such interactions are captured in the Oct4 directly repressing Nanog. In plants, the corresponding reduction of the repression from CLV3 on WUS leads to increased number of stem cells. In particular, induced silencing shows spontaneous reprogramming of meristematic cells [46].

In summary, the need of specific treatment in the mammalian system might be due to the presence of incoherent loops between pluripotency factors. Our simulations show that the removal of incoherence in the ESC system drastically reduces the differences in reprogramming efficiencies in the two kingdoms.

Reprogramming time differences

We monitored the reprogramming time for the ESC (Oct4-Nanog incoherence is present) and SAM model (Fig 3). The reprogramming time distributions show that, at least for high overexpression, the SAM model reprograms faster compared to the ESC model. The reprogramming time distributions obtained from ESC model simulations are more skewed than the ones obtained from SAM model, due to the presence of incoherent loop between Oct4 and Nanog. Also, the variations in reprogramming time are larger for the ESC model.

Altogether, our results suggest that the gene regulatory networks topologies along with the differentiation dynamics are similar for the SAM and ESC systems. However, the reprogramming dynamics differ, the SAM model being more amenable to reprogram from a differentiated state.

Discussion

It appears that simplified single cell models of the mammalian embryonic stem cells and plant stem cells can account for the differences in reprogramming properties, although the network dynamics are similar when it comes to differentiation properties. The 'simple'
reactivation of CLV3/WUS is in nice agreement with plants’ capability to spontaneously create new stem cells in the initialization of new reproductive organs around the main stem [4], as discussed above. This indicates that many of the interactions taking part in the cell state transitions could be simplified to the few components used here in the core regulatory modules. It should be stressed that the Oct-dependent epigenetic effect is included in the ESC model [38]. In contrast such epigenetic mechanism has still not been identified when overexpressing WUS in the SAM [18]. In both kingdoms, over-expression of key genes can be used as a reprogramming recipe [24, 42]. One could also accomplish similar results by suppressing the differentiation gene G. On the other hand since there is very likely more than one differentiation gene in the systems, such an approach might not be sufficient alone, as can be exemplified by the relatively weak phenotypes of the KAN loss of function mutations. Combining over-expression and suppression should lead to a more efficient reprogramming. Indeed, recent experimental results showed that depleting Mbd3, a core member of the Mbd3/NuRD (nucleosome remodelling and deacetylation) repressor complex, together with transcription factors over-expression dramatically improve reprogramming efficiency [47]. Mbd3/NuRD plays also a key role in reprogramming and its increased levels can enhance reprogramming efficiency when co-expressed with reprogramming factor NANOG [48]. The small variations of wiring in the core networks of the two stem cell kingdoms could be due to players, which appear to be specialized, being actually able to hold multiple roles in the mammalian stem cell system e.g. Oct4 [40, 49], Mbd3 [47, 48], while plants are known to use different proteins from the same protein families in regulating differentiation in different tissues [25]. In particular, we found that the incoherent feed

![Fig 3. Reprogramming time distributions for various Oct4 and WUS over-expression levels.](https://doi.org/10.1371/journal.pone.0175251.g003)
forward motif between Oct4 and Nanog could be perturbed, leading to more similar reprogramming behaviour as in plants. For example, a reduction of the repression would lead to more plant like behaviour. In plants, this repression sets the WT regulation of number of stem cells and if this is reduced, spontaneous differentiation appears.

In conclusion, by only studying the dynamics of core modules for stem cell regulation within these simplified SAM and ESC models a clear message emerges— the small variations in wiring in the core regulatory networks can explain why reprogramming is more natural in the plant than the mammalian world. For plants reprogramming is a much-needed competence in contrast to animals where it should be more prohibited.

Methods

The shoot apical meristem model

The plant SAM gene circuit in Fig 1A is based on following interactions: CLV represses WUS [32, 33], WUS positively regulates CLV [5], GSAM (KAN1) and WUS are repressing each other [18].

The corresponding set of equations are obtained for the dynamics of CLV, WUS and GSAM (KAN) with mRNA and protein concentrations denoted by [W], [C], [GSAM] and [w], [c], [gsAM] respectively.

\[
\frac{d[W]}{dt} = \frac{p_0}{1 + p_1[C] + p_2[GSAM]^2} - d_w[W]
\]
\[
\frac{d[w]}{dt} = P_w[W] - d_w[w]
\]
\[
\frac{d[C]}{dt} = \frac{p_3 + p_4[w]}{1 + p_4[w]} - d_c[C]
\]
\[
\frac{d[c]}{dt} = P_c[C] - d_c[c]
\]
\[
\frac{d[GSAM]}{dt} = \frac{p_5}{1 + p_6[w]^2} - d_{GSAM}[GSAM]
\]
\[
\frac{d[gsAM]}{dt} = P_{gsAM}[GSAM] - d_{gsAM}[gsAM]
\]

We use mass action dynamics for protein production and degradation, and a standard Shea-Ackers description for the transcriptional regulation [50], which adds specific constraints on parameters for the underlying elementary reactions for exact description of dynamics and noise distributions [51]. \(P_x\) represents production rates of protein \(x\), \(d_x\) is the degradation rate of molecule \(x\) (mRNA and protein), and the \(p\) parameters relate to the transcriptional regulation. Three examples of parameters sets from our optimizations (see Parameter optimization section) that were used for calculating the reprogramming efficiencies depicted in Fig 2 are shown below in Table 1. It must be noted that we optimized the parameters based on two main constraints: bistability and spontaneous differentiation. For the SAM model we found 43 successful parameter sets, see Table A in S1 File. The protein production rates \(P\) and the mRNA and protein decay rates \(d\) were chosen to be 0.01.
The embryonic stem cell model

The mammalian embryonic stem cell circuit in Fig 1A was built based on following: OCT4 and SOX2 form a complex which induces its own production [34] and also induces and represses NANOG and induces GESC [2, 3, 38], NANOG represses itself [35, 36], GESC (Gata6, Sox17) and NANOG are mutually repressive furthermore GESC represses OCT4/SOX2 [38].

We put forward the following set of ordinary differential equations from a thermodynamic approach [50] describing the behaviour of NANOG, OCT4/SOX2 and the differentiation gene GESC (GATA6; SOX17) with mRNA and protein concentrations denoted by [N], [OS], [GESC] and [n], [os], [gESC] respectively.

\[
\frac{d[N]}{dt} = \frac{p_1[os](p_0 + p_1[os])}{1 + p_1[os][p_1[os] + p_2[n] + p_4[os]] + p_1[os][gESC]^2} - d_{n}[N]
\]

\[
\frac{d[n]}{dt} = P_n[N] - d_{n}[n]
\]

\[
\frac{d[OS]}{dt} = \frac{p_5 + p_6[os]^2}{1 + p_6[os] + p_7[gESC]^2} - d_{os}[OS]
\]

\[
\frac{d[os]}{dt} = P_{os}[OS] - d_{os}[os]
\]

\[
\frac{d[GESC]}{dt} = \frac{p_8 + p_9[gESC]^2 + p_{10}[os]}{1 + p_9[gESC]^2 + p_{11}[n]^2} - d_{GESC}[GESC]
\]

\[
\frac{d[gESC]}{dt} = P_{gESC}[GESC] - d_{gESC}[gESC]
\]

Parameters are defined as for the SAM model. Three examples of parameters sets from our optimizations (see Parameter optimization section below) used for the calculations of reprogramming efficiencies depicted in Fig 2 are shown in Table 2.

Table 1. Three SAM parameters sets examples out of 43 parameter sets that were optimized based on bistability and spontaneous differentiation constraints.

| Parameters | $p_0$ | $p_1$ | $p_2$ | $p_3$ | $p_4$ | $p_5$ | $p_6$ | $p_7$ | $p_8$ | $p_9$ | $p_{10}$ | $p_{11}$ |
|------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|--------|--------|
| Set 1      | 0.11  | 0.05  | 5.58  | 0.04  | 0.08  | 0.84  | 0.002 |       |       |       |        |        |
| Set 2      | 0.04  | 0.02  | 3.63  | 0.003 | 1.90  | 1.03  | 0.02  |       |       |       |        |        |
| Set 3      | 0.05  | 0.74  | 4.57  | 0.03  | 0.49  | 0.78  | 0.01  |       |       |       |        |        |

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Table 2. Three ESC parameters sets examples out of 25 parameter sets that were optimized based on bistability and spontaneous differentiation constraints.

| Parameters | $p_0$ | $p_1$ | $p_2$ | $p_3$ | $p_4$ | $p_5$ | $p_6$ | $p_7$ | $p_8$ | $p_9$ | $p_{10}$ | $p_{11}$ |
|------------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|--------|--------|
| Set 1      | 50    | 1.50  | 0.40  | 0.90  | 0.40  | 0.01  | 1.00  | 1.00  | 0.005 | 0.05  | 1.20   | 0.35   |
| Set 2      | 50    | 1.60  | 0.60  | 0.90  | 0.32  | 0.007 | 1.34  | 1.00  | 0.005 | 0.04  | 1.55   | 0.35   |
| Set 3      | 50    | 1.71  | 0.42  | 0.90  | 0.30  | 0.006 | 0.38  | 0.30  | 0.005 | 0.08  | 1.20   | 0.35   |

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are not local. For the ESC model 25 parameter sets accounted for bistability and spontaneous differentiation in the system see Table B in S1 File. The protein production rates $P$ and the mRNA and protein decay rates $d$ were considered to be 0.01 for all species.

**Differentiation simulations**

For both SAM and ESC gene regulatory networks we developed computational models consisting of ordinary differential equations shown in the sections above. The Shea- Ackers approach [50] was used for describing transcription while mass action kinetics were employed for describing mRNA and protein degradation along with production of protein. We consider the initial conditions of the system to be the state where all the genes have high expression except $G_{\text{SAM}}$ and $G_{\text{ESC}}$ (stem cell state). To get familiar with the models and their normal behaviour several stochastic simulations were conducted [52], two examples corresponding to two single cells are presented in Fig 1B. We also conducted multiple simulations corresponding to cells colonies. Fig A in S1 File shows WUS and OCT4-SOX2 gene expression distribution at a time point towards the end of the simulations. The distributions demonstrate the robust bistability of the SAM and ESC systems. The parameters used for producing the results shown in Fig 1B and Fig A in S1 File are presented in the first line of Tables 1 and 2. Please note that $p_3$ in the ESC model takes a value of 1.5 for these simulations. The spontaneous differentiation time for both models was investigated. The differentiation time was taken to be time interval between start of simulation and the moment that both the types of mRNA, characteristic for the stem cell state (e.g. OCT4/SOX2 and NANOG for ESC), had lower levels than the differentiation promoting genes. Before saving the differentiation time, the simulation ran for yet $10^4$ reactions to separate short fluctuations from real state shifts. If the system remained in the differentiated state for these $10^4$ reactions we consider the simulation to be a successful simulation of the differentiation process.

We conducted 100 stochastic simulations for each parameter set, fixing a maximum number of reactions equal to $10^7$, as a stopping criterion. Matlab (The Mathworks) was used to solve the differential equations and conduct stochastic Gillespie simulations.

**Reprogramming simulations**

We initiate our simulations by considering the systems to be in a state where $G_{\text{SAM}}$ and $G_{\text{ESC}}$ are at high values (differentiated state) while the rest of the genes (the stem cell specific ones) are at low values. The promotion of stem cell specific genes was implemented by adding a constant production (reprogramming force) to the equation for OCT4/SOX2 mRNA rate of change in the ESC model and correspondingly for WUS in the SAM model. If the addition of this constant resulted in a switch of the system to a state where $G_{\text{SAM}}$ and $G_{\text{ESC}}$ expression are low while the other genes expression are high then we considered the reprogramming to be successful. The reprogramming force of over-expression level for OCT4/SOX2 and WUS were varied in a logarithmic fashion. For each value of over-expression level, simulation data was collected to measure the degree of success (the efficiency) for the reprogramming method i.e. the time required to reprogram the cell and the fraction of reprogrammed cells. Initially, 100 simulations were performed for each parameter set and each value of reprogramming force with the stop criterion of $10^6$ reactions. In order to gather statistics 400 additional simulations were made for the reprogramming force shown in Fig 2.

**Parameter optimization**

To optimize the parameter sets for the SAM and ESC models we used the simulated annealing global optimization algorithm [53, 54]. We optimized the parameters based on the constraints
that the systems must be bistable and spontaneous transition from pluripotent state to differentiated state must occur during stochastic simulations. To take into account the bistability of the models two cost functions were needed, one for the stem cell state and another one for the differentiated state. The stem cell state is defined as in [38] as a state with high levels of WUS, CLV3, OCT4/SOX2, NANOG and low levels of G_{SAM} and G_{ESC} and vice versa for the differentiated state. We use the following cost function:

\[
\begin{align*}
  f_{\text{stem}} &= \frac{1}{N} \sqrt{\sum_{i=1}^{N} (x_{\text{stem}}^{i,\text{model}} - x_{\text{stem}}^{i,\text{opt}})^2} \\
  f_{\text{diff}} &= \frac{1}{N} \sqrt{\sum_{i=1}^{N} (x_{\text{diff}}^{i,\text{model}} - x_{\text{diff}}^{i,\text{opt}})^2} \\
  f_i &= f_{\text{stem}} + f_{\text{diff}}
\end{align*}
\]

where \( N \) is the number of different molecules (\( N = 3 \)), \( x_{\text{stem}}^{i,\text{model}} \) is the mRNA concentration of the species type \( i \) (e.g. SAM \( i = \{W, C, G_{SAM}\} \)) obtained from the model simulation with initial conditions in the stem cell state, while the \( x_{\text{diff}}^{i,\text{model}} \) is obtained from model simulation starting in differentiated state, \( x_{\text{stem}}^{i,\text{opt}} \) is the optimal value of the concentration of mRNA of species type \( i \).

Although the above cost function specifies the bistability condition a third cost function was introduced for taking into account the differences between stem cell and differentiated cell states:

\[
\begin{align*}
  f_2 &= \left| \frac{1}{N} \sqrt{\sum_{i=1}^{N} (x_{\text{stem}}^{i,\text{opt}} - x_{\text{stem}}^{i,\text{model}})^2} - \frac{1}{N} \sqrt{\sum_{i=1}^{N} (x_{\text{diff}}^{i,\text{opt}} - x_{\text{diff}}^{i,\text{model}})^2} \right|
\end{align*}
\]

The second cost function (\( f_2 \)) penalized solutions where only one of the states was fitted well. Furthermore it facilitated to distinguish between successful and unsuccessful optimizations. \( x_{\text{stem}}^{i,\text{model}} \) was obtained from conducting model simulations using a fourth order Runge-Kutta with adaptive step size method. \( x_{\text{stem}}^{i,\text{opt}} \) values resemble experimental observations with high values varying between 80 and 120 and the low value between 0 and 20.

We used an annealing schedule similar to the one in [54]. We define the acceptance ratio as:

\[
X(c) = \frac{\text{number of accepted transitions}}{\text{number of proposed transitions}}
\]

where \( c \) is the control parameter \( c_{\text{new}} = k_{\text{cool}} c_{\text{old}} \) with \( k_{\text{cool}} = 0.99 \). The parameter configurations can be viewed as states in a Markov chain. The length of each Markov chain should be large enough for the algorithm to be able to explore the cost function landscape around the point defined by the configuration obtained in the previous step in the cooling process. The higher dimension a problem has, the longer the chain should be. This feature is incorporated in the following expression for the length: \( L = L_0 n \), with standard length \( L_0 = 10 \) and \( n \) being the dimension of the problem.

**Supporting information**

S1 File. Contains Table A with all optimized parameters for the SAM model, Table B with all optimized parameters for ESC model, Table C showing sensitivity analysis results and
Fig A showing distribution results from population level stochastic simulations.

(PDF)

**Author Contributions**

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**Supervision:** HJ CP.

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**Visualization:** VO JN.

**Writing – original draft:** VO HJ CP.

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**References**

1. Nichols J, Zevnik B, Anastassiadis K, Niwa H, Klewe-Nebenius D, Chambers I, et al. Formation of pluripotent stem cells in the mammalian embryo depends on the POU transcription factor Oct4. Cell. 1998; 95: 379–391. PMID: 9814708

2. Mitsui K, Tokuzawa Y, Itoh H, Segawa K, Murakami M, Takahashi K, et al. The homeoprotein Nanog is required for maintenance of pluripotency in mouse epiblast and ES cells. Cell. 2003; 113: 631–642. PMID: 12787504

3. Boyer LA, Lee TI, Cole MF, Johnstone SE, Levine SS, Zucker JP, et al. Core transcriptional regulatory circuitry in human embryonic stem cells. Cell. 2005; 122: 947–956. https://doi.org/10.1016/j.cell.2005.08.020 PMID: 16153702

4. Mayer KFX, Schoof H, Haecker A, Lenhard M, Jürgens G, Laux T. Role of WUSCHEL in Regulating Stem Cell Fate in the Arabidopsis Shoot Meristem. Cell. 1998; 95: 805–815. PMID: 9865698

5. Fletcher JC, Brand U, Running MP, Simon R, Meyerowitz EM. Signaling of cell fate decisions by CLAVATA3 in Arabidopsis shoot meristems. Science. 1999; 283: 1911–1914. PMID: 10082464

6. Clark SE, Williams RW, Meyerowitz EM. The CLAVATA1 gene encodes a putative receptor kinase that controls shoot and floral meristem size in Arabidopsis. Cell. Elsevier; 1997; 89: 575–585.

7. Lee D-S, Shin J-Y, Tonge PD, Puri MC, Lee S, Park H, et al. An epigenomic roadmap to induced pluripotency reveals DNA methylation as a reprogramming modulator. Nat Commun. 2014; 5: 5619. https://doi.org/10.1038/ncomms6619 PMID: 25493341

8. Hussein SMI, Puri MC, Tonge PD, Benevento M, Corso AJ, Clancy JL, et al. Genome-wide characterization of the routes to pluripotency. Nature. 2014; 516: 198–206. https://doi.org/10.1038/nature14046 PMID: 25503233

9. Loh Y-H, Zhang W, Chen X, George J, Ng H-H, JmjD1a and JmjD2c histone H3 Lys 9 demethylases regulate self-renewal in embryonic stem cells. Genes Dev. 2007; 21: 2545–2557. https://doi.org/10.1101/gad.1588207 PMID: 17938240
10. Soufi A, Donahue G, Zaret KS. Facilitators and impediments of the pluripotency reprogramming factors’ initial engagement with the genome. Cell. 2012; 151: 994–1004. https://doi.org/10.1016/j.cell.2012.09.045 PMID: 23159369

11. Chen J, Liu H, Liu J, Qi J, Wei B, Yang J, et al. H3K9 methylation is a barrier during somatic cell reprogramming into iPSCs. Nat Genet. 2013; 45: 34–42. https://doi.org/10.1038/ng.2491 PMID: 23202127

12. Silva J, Nichols J, Theunissen TW, Guo G, van Oosten AL, Barrandon O, et al. Nanog is the gateway to the pluripotent ground state. Cell. 2009; 138: 722–737. https://doi.org/10.1016/j.cell.2009.07.039 PMID: 19703398

13. Costa Y, et al. NANOG-dependent function of TET1 and TET2 in establishment of pluripotency. Nature. 2013; 495:370–374. https://doi.org/10.1038/nature11925 PMID: 23395962

14. Olariu V, Lövkvist C, Sneppen K. Nanog, Oct4 and Tet1 interplay in establishment pluripotency. Scientific Reports 2016; 6:

15. Schoof H, et al. The Stem Cell Population of Arabidopsis Shoot Meristems Is Maintained by a Regulatory Loop between the CLAVATA4 and WUSCHEL Genes. Cell. 2000; 100(6):635–644. PMID: 10761929

16. Yadav RK, et al. WUSCHEL protein movement mediates stem cell homeostasis in the Arabidopsis shoot apex. Genes Dev. 2011; 25:2025–2030. https://doi.org/10.1101/gad.17258511 PMID: 21979915

17. Daum G, et al. A mechanistic framework for noncell autonomous stem cell induction in Arabidopsis. Proc Natl Acad Sci U S A. 2014; 111(40):14619–24. https://doi.org/10.1073/pnas.1406446111 PMID: 25245676

18. Yadav RK, Perales M, Gruel J, Ohno C, Heisler M, Girke T, et al. Plant stem cell maintenance involves direct transcriptional repression of differentiation program. Mol Syst Biol. 2013; 9: 654. https://doi.org/10.1038/msb.2013.8 PMID: 23549482

19. Sun B, Ito T. Regulation of floral stem cell termination in Arabidopsis. Front Plant Sci. Frontiers; 2015; 6: 17.

20. Sun B, Locic R, Guo S, He Z, Gan E-S, Huang J, et al. Timing Mechanism Dependent on Cell Division Is Invoked by Polycomb Eviction in Plant Stem Cells. Science. American Association for the Advancement of Science; 2014; 343: 1248559.

21. Kornet N, Scheres B. Stem Cell Factors in Plants: Chromatin Connections. Cold Spring Harb Symp Quant Biol. Cold Spring Harbor Laboratory Press; 2008; sqb.2008.73.043.

22. Pikaard CS, Mittelsten Scheid O. Epigenetic regulation in plants. Cold Spring Harb Perspect Biol. 2014; 1:6(12)

23. Skoog F, Miller CO. Chemical regulation of growth and organ formation in plant tissues cultured in vitro. Symp Soc Exp Biol. 1957; 11: 118–130. PMID: 13486467

24. Takahashi K, Yamakawa S. Induction of Pluripotent Stem Cells from Mouse Embryonic and Adult Fibroblast Cultures by Defined Factors. Cell. 2006; 126: 663–676. https://doi.org/10.1016/j.cell.2006.07.024 PMID: 16904174

25. Heidstra R, Sabatini S. Plant and animal stem cells: similar yet different. Nat Rev Mol Cell Biol. 2014; 15: 301–12. https://doi.org/10.1038/nrm3790 PMID: 24755932

26. Faunes F, Hayward P, Descalzo SM, Chatterjee SS, Balayo T, Trotta J, et al. A membrane-associated β-catenin/Oct4 complex correlates with ground-state pluripotency in mouse embryonic stem cells. Development. The Company of Biologists Limited; 2013; 140: 1171–1183.

27. Zhao Z, et al. Hormonal control of the shoot stem-cell niche. Nature. 2010; 465:1089–1092. https://doi.org/10.1038/nature09126 PMID: 20577215

28. Su YH, Liu YB, Zhang XS. Auxin–Cytokinin Interaction Regulates Meristem Development. Molecular Plant. 2011; 4(4):616–620. https://doi.org/10.1093/mp/ssr007 PMID: 21357646

29. Clark SE, Running MP, and Meyerowitz EM. Clavata3 is a speciﬁc regulator of shoot and floral meristem development affecting the same processes as clavata1, Development, 1995; 121: 2057–2067.

30. Kayes JM, Clark SE. Clavata2, a regulator of meristem and organ development in Arabidopsis. Development, 1998; 125:3843–3851. PMID: 9729492

31. Laux T, Mayer KFX, Berger J, Jurgens G. The wuschel gene is required for shoot and floral meristem integrity in arabiadopsis,. Development, 1996; 122:87–96. PMID: 8565856

32. Brand U, Fletcher JC, Hobe M, Meyerowitz EM, Simon R. Dependence of stem cell fate in Arabidopsis on a feedback loop regulated by clv3 activity, Science 2000; 289: 617–619. PMID: 10915624

33. Lenhard HSM, Haeker A, Mayer KFX, Jurgens G. The stem cell population of Arabidopsis shoot meristems is maintained by a regulatory loop between the clavata and wuschel genes, Cell 2000; 100:635–644. PMID: 10761929
34. Chew JL, Loh YH, Zhang WX, Chen W, Tam WL, Yeap LS, et al. Reciprocal transcriptional regulation of pou5f1 and sox2 via the oct4/sox2 complex in embryonic stem cells. Molecular and Cellular Biology 2005; 25(4):6031–6046.

35. Navarro P, Festuccia N, Colby D, Gagliardi A, Mullin NP, Zhang W, et al. Oct4/sox2-independent nanog autorepression modulates heterogeneous nanog gene expression in mouse es cells. The EMBO Journal, 2012; 31:4547–4562. https://doi.org/10.1038/emboj.2012.321 PMID: 2317892

36. Fidalgo M, Faiola F, Pereira CF, Saunders JDA, Gingold J, Schaniel C et al. Zfp281 mediates nanog autorepression through recruitment of the nuc complex and inhibits somatic cell reprogramming. Proceedings of the National Academy of Sciences of the United States of America, 2012; 109:16202–16207. https://doi.org/10.1073/pnas.1208533109 PMID: 22988117

37. Silva J, Smith A. Capturing pluripotency. Cell, 2008; 132:532–536. https://doi.org/10.1016/j.cell.2008.02.006 PMID: 18295569

38. Chickarmane V, Olariu V, Peterson C. Probing the role of stochasticity in a model of the embryonic stem cell: heterogeneous gene expression and reprogramming efficiency. BMC Syst Biol, BioMed Central Ltd; 2012; 6: 98.

39. Papapetrou EP, Tomishima MJ, Chambers SM, Mica Y, Reed E, Menon J, et al. Stoichiometric and temporal requirements of Oct4, Sox2, Klf4, and c-Myc expression for efficient human iPSC induction and differentiation. Proc Natl Acad Sci. National Acad Sciences; 2009; 106: 12759–12764.

40. Livigni A, Peradze H, Sharov AA, Chia G, Hammachi F, Migueles RP, et al. A conserved Oct4/POU V-dependent network links adhesion and migration to progenitor maintenance. Curr Biol. 2013; 23: 2233–2244. https://doi.org/10.1016/j.cub.2013.09.048 PMID: 24210613

41. Bouchabeké-Coussa O, et al. Wuschel overexpression promotes somatic embryogenesis and induces organogenesis in cotton (Gossypium hirsutum L.) tissues cultured in vitro. Plant Cell Rep. 2013; 32(5):675–86. https://doi.org/10.1007/s00299-013-1402-9 PMID: 23543366

42. Gallois J-L, Woodward C, Reddy VG, Sablowski R. Combined SHOOT MERIS TEMLESS and WUSCHEL trigger ectopic organogenesis in Arabidopsis. The Company of Biologists Limited; 2002; 129: 3207–3217.

43. Kim J. B. et al. Oct4-induced pluripotency in adult neural stem cells. Cell. 136, 411–419 (2009). https://doi.org/10.1016/j.cell.2009.01.023 PMID: 19203577

44. Tsai S. Y. et al. Single transcription factor reprogramming of hair follicle dermal papilla cells to induced pluripotent stem cells. Stem Cells 29, 964–971 (2011). https://doi.org/10.1002/stem.649 PMID: 21563278

45. Wu T. et al. Reprogramming of trophoblast stem cells into pluripotent stem cells by Oct4. Stem Cells. 29, 755–763 (2011). https://doi.org/10.1002/stem.617 PMID: 21305674

46. Reddy GV, Meyerowitiz EM. Stem-cell homeostasis and growth dynamics can be uncoupled in the Arabidopsis shoot apex. Science. 2005; 310(5748):663–7. https://doi.org/10.1126/science.1116261 PMID: 16210497

47. Rais Y, Zviran A, Geula S, Gafni O, Chomsky E, Viukov S, et al. Deterministic direct reprogramming of somatic cells to pluripotency. Nature. 2013; 502: 65–70. https://doi.org/10.1038/nature12587 PMID: 24048479

48. Dos Santos RL, Tosti L, Radzisheuska A, Caballero IM, Kaji K, Hendrich B, et al. MBD3/NuRD facilitates induction of pluripotency in a context-dependent manner. Cell Stem Cell. 2014; 15: 102–110. https://doi.org/10.1016/j.stem.2014.04.019 PMID: 24835571

49. Radzisheuska A, Chia GL, Bin, dos Santos RL, Theunissen TW, Castro LFC, Nichols J, et al. A defined Oct4 level governs cell state transitions of pluripotency entry and differentiation into all embryonic lineages. Nat Cell Biol. 2013; 15: 579–590. https://doi.org/10.1038/ncb2742 PMID: 23629142

50. Shea MA, Ackers GK. The OR control system of bacterio phage lambda a physical-chemical model for gene regulation. Journal of Molecular Biology, 1985; 181(2):211–230. PMID: 3157005

51. Thomas P. et al. Communication: limitations of the stochastic quasi-steady-state approximation in open biochemical reaction networks. J Chem Phys, 2011; 135:181103–181103. https://doi.org/10.1063/1.3661156 PMID: 22088045

52. Gillespie DT. Stochastic simulation of chemical kinetics., Annual Review of Physical Chemistry 2007; 58:35–55. https://doi.org/10.1146/annurev.physchem.58.032806.104637 PMID: 17037977

53. Kirkpatrick S, Gelatt CD, Vecchi MP. Optimization by simulated annealing. Science, 1983; 220 (4598):671, 680, 1983. https://doi.org/10.1126/science.220.4598.671 PMID: 17813860

54. Cerny V. Thermodynamical approach to the traveling salesman problem: An efficient simulation algo- rithm. Journal of optimization theory and application. 1985; 45(1):41–51.