Versioning biological cells for trustworthy cell engineering

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“Full-stack” biotechnology platforms for cell line (re)programming are on the horizon, thanks mostly to (a) advances in gene synthesis and editing techniques as well as (b) the growing integration of life science research with informatics, the internet of things and automation. These emerging platforms will accelerate the production and consumption of biological products. Hence, traceability, transparency, and—ultimately—trustworthiness is required from cradle to grave for engineered cell lines and their engineering processes. Here we report a cloud-based version control system for biotechnology that (a) keeps track and organizes the digital data produced during cell engineering and (b) molecularly links that data to the associated living samples. Barcoding protocols, based on standard genetic engineering methods, to molecularly link to the cloud-based version control system six species, including gram-negative and gram-positive bacteria as well as eukaryote cells, are shown. We argue that version control for cell engineering marks a significant step toward more open, reproducible, easier to trace and share, and more trustworthy engineering biology.
Engineering biology is exploding with advances ranging from new genome editing tools\(^1\), to genetically encodable materials for advanced sensing of cells physiological states, electrical fields and mechanical stresses\(^2,3\), programmable and functional microbial-based living materials\(^4\), environmental remediation and pollution control\(^5\) to advanced in vivo data storage\(^6\). Moreover, these advances in fundamental science are rapidly translating into new companies\(^7\) and consumer products\(^8\), which within the first half of this century, are set to impact most areas of our lives.

Perhaps the most convincing example of the pace of progress is the global scientific response to the current SARS-CoV-2 pandemic. In a matter of weeks after detecting the outbreak, the virus was isolated, its genome sequenced and published\(^9\) and made available for research. Slightly less than a year later multiple vaccines were already being deployed to combat the virus. This would have been unimaginable just 10 years ago. Although this is an extreme example arising out of an extreme situation, it is to be expected that with the commoditization of synthetic DNA and the wider availability of powerful gene-editing tools\(^10\), the number of engineered strains will rapidly increase. Indeed, cheaper DNA synthesis technology and the development of high throughput, automated, cloning processes allows the creation of large plasmid and combinatorial DNA libraries\(^11,12\) in a matter of days, including the modification of recalcitrant species’ genomes\(^13\), which previously were difficult to edit.

And yet, while engineering biology has changed profoundly in the last few years, there are still deep gaps in the way the process of strain engineering is done and disseminated. For example, engineering a “synthetic biology agent”\(^14\) produces large quantities of information: published articles, protocols, notebooks, models, databases, sequencing and other types of data (e.g., metabolomics, proteomics, lipomics, etc.). Combined, all these information sources may add up to terabytes of data but only a relatively small percentage of it is being made available when results are published in specialized outlets. This gap in scientific practice has led to an ongoing crisis in cell line misidentification\(^15,16,17\), a recognized lack of reproducibility\(^18\), sometimes causing high profile retractions\(^19\) and often resulting in weakening public attitudes to new and emerging technologies.

It is thus clear that this gap in scientific practice requires a response on multiple fronts, to which this paper contributes in a number of practical ways with the introduction of CellRepo as a community resource.

CellRepo is a version control system for cell engineering. Version control is the practice of monitoring modifications in the source code of computer programs or other (digital) objects, which is assisted by the use of special software that keeps track of the code, its changes over time, the changes’ authors and other metadata. CellRepo integrates, on the one hand, a cloud-based version control software for tracking cell lines’ digital footprints and, on the other hand, living samples’ molecular barcoding protocols to link the biological sample back to their cloud digital information. CellRepo relies on two fundamental pillars. Firstly, during the process of engineering a new strain, changes introduced to the cell line are recorded in “commits” that include information such as the genotype, phenotype, author of the modifications, laboratory protocols, characterization profiles, etc. (Fig. 1a). The history of commits tracks the digital footprint produced during the process of cell engineering. The second pillar is the physical linking of a living sample to a commit via the chromosomal introduction of a unique barcode related to the commit (Fig. 1b).

Genomic barcodes have been recently recommended as the way forward for tagging\(^18,20\) synthetic biology chassis with unique identifiers for the sake of traceability, intellectual property issues and environmental risk assessment (ERA)\(^21\). Current schemes for curbing the propagation of genetically modified microorganisms with genetic firewalls or conditional killing systems are still insufficient to guarantee certainty of containment\(^22\). In this context, barcodes appear as either an alternative or as a complement to such firewalls for a sound ERA of agents designed for deliberate release—or accidentally escaped thereof. Genomically inserted identifiers instantly refer to digital twins with all available documentation on the live construct at stake (see below), not only in terms of species and genetic pedigree but also regarding safety aspects and indications for countermeasures in case of undesirable propagation. In that sense, barcodes may ease the current emphasis on containment toward a more realistic scenario of management\(^23\), thereby facilitating the regulatory and approval process\(^14,21\).

To demonstrate the wide applicability of CellRepo, we show how the platform—in conjunction with well-established peer-reviewed protocols to genetically engineer various organisms—can be applied to six of the most important and diverse microbial species used in both academia and industry (Escherichia coli, Bacillus subtilis, Streptomyces albusdoflavus, Pseudomonas putida, Saccharomyces cerevisiae and Komagataella phaffii—previously known as Pichia pastoris). It is expected that the same principles can be applied to more, if not all, species which have been already domesticated and engineered.

**Results**

**CellRepo is a cloud version control for engineering biology.** We created a cloud-based community resource built on top of a modern software engineering stack for web applications. As in any cloud-based application, the user needs to register, providing a name, e-mail address and password. An avatar picture can be uploaded to personalize the experience and to be more recognizable by other users (e.g., collaborators). After registering, the user will receive a confirmation by e-mail. Finally, the user can sign in by typing the registration e-mail and password. Once a user signs in, they land on the homepage (Fig. 2a) that contains everything needed to build repositories of engineered strains, manage their accounts and the teams they work with. From the initial page, it is also possible to access the system documentation (“Knowledge Base”). The blue upper horizontal quick menu links to all the aforementioned features and is present on every page on the website. This menu also contains a search bar. This allows users to look for repositories and commits accessible to them (i.e., cell repositories they own, that belong to their teams or public cell repositories) and look for identifiers to find the documentation on specific strains/plasmids.

The first step to start a repository is to select a species. The server is linked to up-to-date databases of organisms (Fig. 2b). This ensures that the users are always able to use the species they need and that these are well documented. To ease the finding of new species to work with, the users need to pre-select them from the database and add the species to their unique list of in-use organisms.

Repositories are projects or experiments (e.g., compound production, protein expression, etc.) and are usually linked to a specific species. Metadata information like the name and description of the project, as well as information about the purpose or how to use a specific strain repository, can be added. Repositories may have different “visibilities”: public (anyone can see the content of the repository), team (visible just for members of the same team or laboratory) and private (just the user can see and add changes). A user may change its repository visibility at any point in time. A repository can have many branches and in turn, branches are made of commits. The name of the initial branch can be set during the creation of the repository.
A repository (Fig. 3a) has a main or leader branch (named during the repository creation) and many other branches. Each branch represents a new direction or idea the users want to pursue in their cell engineering activities (e.g., novel protocols, different gene edition order, etc.).

A commit (Fig. 3b) captures the status of the engineered strain at a specific point in time. The amount of information contained in a commit is up to the user (it can be as simple as a new strain name or as complex as a brand-new strain creation by modifying the genome and adding several documents). In addition, the commits are the containers of the uploaded documentation (which can be in the form of documents, models, sequences, etc.).

Once in a repository, the user can choose a branch to commit. The “new commit” button opens a form in which various types of information can be inserted. For example, the user can name and describe the commit (what is being done? why? what for?). Importantly, all types of documentation (supporting the commit) can be uploaded on this page such as construct sequence,
Electrophoresis gel pictures, SBOL files, growths and fluorescence curves, sequencing results, automation worklist instructions, computer models, etc. The user can also provide genotype and phenotype information, the storage location of the strain, safety information, acceptable material transfer agreements for the strain, etc.

The user can choose the level of granularity of commits that best fits its laboratory practice, e.g., a commit might represent a single cell modification or multiple multi-loci genetic changes.

When creating a new commit, the user can decide whether the change is important enough (e.g., a milestone) to be physically barcoded into the cell. If that is the case, the system allows the generation of a unique barcode sequence. The barcode can then be synthesized and inserted into the strain. Once created, the commit will be linked unequivocally to the strain carrying the barcode sequence.

CellRepo allows users to be part of collaboration “teams” for cell engineering. Team members of a strain repository can make commits and create new branches to the cell line history. Furthermore, teams of researchers can share repositories, track strains and be up to date on the experiments being carried out in their projects. Creating a new team is as easy as providing a name to the team and adding CellRepo users to it. Once established, it is possible to see all the members and the shared repositories and keep track of the activity taking place on the repository.

In vivo barcoding experiments. Different barcoding protocols (detailed step-by-step protocols can be found in the extensive Supplementary material) were assessed for E. coli, B. subtilis, S. albidoslavaus, P. putida, S. cerevisia and K. phaffii—previously known as P. pastoris. These protocols are used to introduce into the chromosome of the cell the barcodes that are automatically generated by CellRepo when a user creates a new commit in the version control system. CellRepo maps the unique commit identifier into DNA sequences that are then used as barcodes. All the tested barcoding procedures successfully barcoded the target species (Supplementary Table 5). URL links and QR codes for all...
CellRepo repositories for these experiments can be found in Supplementary Table 6.

For all species tested, barcodes are genetically, physiologically innocuous and stable over a range of growth conditions. Barcoding a strain should have little to no effect on its growth profile; growth profiles of barcoded cells were compared to wild-type (i.e., non-barcoded) strains (Supplementary Fig. 5).

The six growth profiles show no significant differences between barcoded strains and the corresponding non-barcoded parental strains. This confirms that the barcode insertion has little effect on the growth of the different species.

We also evaluated whether or not the barcoding protocols introduced unplanned mutations in the recipient cell. For instance, this can help choose a specific barcoding method over another. To do this, the whole genome of the barcoded strains and the wild-type strains used was sequenced. The results can be found in Supplementary Tables 7–12.

E. coli results show that the three clones barcoded using Lambda-Red method had the same point mutation in an intergenic region (Supplementary Table 7). This may be explained by the fact that the initial colony chosen to start the insertion process already contained the mutation or it was acquired during the process. In any case, the mutation is intergenic and does not seem to affect the cells. In one of the clones barcoded using gRNA1, two-point mutations appear in different CDS. Strains barcoded using gRNA2 do not show any mutation.

B. subtilis was barcoded using three different methods. Both CRISPR (only one gRNA tested) and Toxin-mediated barcoded cells show no mutations in all the different clones. In one of the barcoded strains using Cre-Lox, a mutation appears. In a different clone, two different SNPs could be detected. All the mutations are in different CDS (Supplementary Table 8).

For P. putida, two clones were barcoded using the CRISPR/targetron system and both showed one mutation in different CDS (Supplementary Table 9).

S. cerevisiae was barcoded using two methods. In both of them, most of the mutations that appear are tandem repeat related. These could be acquired during the insertion process or could be sequencing artifacts related to this type of repetitive sequence. Two strains barcoded using Cre-Lox show single point mutations. Four mutations (tandem repeats) are observed in Strain 2 (CRISPR). A single point SNP can be observed in Strain 3 (Supplementary Table 10).

One of the two strains of K. phaffii shows two-point mutations in intergenic regions (Supplementary Table 11).

Finally, S. albidosflavus was barcoded using CRISPR. NGS analysis of S. albidosflavus shows a larger number of variants. Strain 1 barcoded with gRNA1 shows two different base-pair changes in different CDS. However, no mutations were found for Strain 2 (gRNA1) (Supplementary Table 12). The three strains produced
using gRNA2 count six, three and two SNPs, respectively. In eukaryotes, CRISPR-caused double-strand breaks (DSB) can be repaired by non-homologous end joining (NHEJ) or homologous recombination (HR) in the presence of a repair template. NHEJ repair is usually imprecise and indels occur. In the case of S. cerevisiae, however, it has been observed that NHEJ hugely decreases cell survival and, when a repair template is provided, HR is the prevalent repair mechanism in this species. Prokaryotes, on the other hand, usually lack NHEJ repair mechanisms. Nevertheless, it has been described in Streptomyces coelicolor among other bacterial species. In this Actinobacteria, closely related to S. albidoflavus, researchers knocked out genes using CRISPR without providing a repair template and allowing the native NHEJ system to—wrongly—repair the DSB. It may be possible that the gRNAs caused CRISPR off-target activity that was repaired by NHEJ causing mutations to appear. Together with the fact that the genome of S. albidoflavus has a high GC content and produced the worst sequencing quality of the analyzed species, this may explain the higher mutation count. Even though other mechanisms may explain the observed variation (see next), users wanting to use pSA-CRISPR-gRNA2 should bear in mind these extra mutations.

The NGS analysis shows that for all the sequenced strains the number of total mutations found in each method is low. The mutations are not constant in the different replicates sent to sequencing. We hypothesize that the mutations (if not sequencing artifacts) are caused by the natural mutation rate in each species during several cycles of growth (both in liquid and solid media). This is supported by. In this S. coelicolor CRISPR edition experiment, the control strain, in which an empty (no target) gRNA was provided to the cells caused a total of seven mutations, three of which were in coding regions. Similar results were found in a CRISPR experiment in S. cerevisiae where they detected 10 SNPs that were probably caused by the successive transformation rounds required for the experiment.

Importantly, we found no structural variants in any of the sequenced strains. The NGS analysis suggests that the barcoding procedures do not change the genome of the strains more than what would be expected while carrying out conventional genetic engineering protocols. CellRepo users can use this information to choose the barcoding method specific for each species that best fits them.

We also carried out stability evaluation of the barcodes where the stability of the barcode sequences was assessed under five different growth conditions. The barcodes were stable both in terms of presence (Supplementary Table 13) and sequence integrity after the long-term experiments (Supplementary Figs. 6–35). Finally, the usage of barcoded strains as a way to track the dissemination of GMOs is described (Supplementary Fig. 36). In the particular case of a gene drive (which has been proposed as a solution to some infectious diseases transmitted to humans from animal and insect vectors), barcode sequences could pinpoint the source of the released modified organisms (29) (intentionally or accidentally) in the environment.

To assess that the barcode stays in the insertion site and that its sequence is still retrievable even after long periods of growth, we ran on all six species five different experiments for 10 days.

As a preliminary experiment, for each condition, the final day single colonies were restreaked and the barcode region was PCR amplified and sent to Sanger sequencing (Eurofins Genomics). For all the colonies tested, we were able to confirm the barcode presence by PCR in all the cases. Supplementary Table 13 describes in detail the sequencing results of this experiment.

To have a more thorough view of what happened to the barcode sequence during the long-term experiments, an NGS analysis of the PCR purified product of the barcoded region of the cell population in conditions 1–4 was performed.

Mutation analysis of the barcode sequences for all species can be found in Supplementary Figs. 6–35. The percentage of reads showing either INDELs or base-pair changes stayed at the same value as the one observed in the control experiment (lower than 0.05%).

Both the single colonies and the population level experiments show that the barcode was still present after the long-term incubation period and that the sequence was stable on all five experimental conditions.

**Barcodes provide a backtrack signal for GMO dispersion experiments.** Gene drive technology allows the researchers to propagate a specific genetic modification through a population. The scientific community needs to assess the risk of this kind of research. Barcode sequences can be helpful in this matter and uniquely identify the laboratories where a gene drive experiment was carried out, the purpose of the modification and any other relevant data (e.g., safety measures implemented).

Supplementary Fig. 536a graphically describes the gene drive molecular mechanism. The barcode identifier was coupled with the intended modification (ADE2 deletion cassette). Supplementary Fig. 36b shows that the barcoded cells (red pigment) can survive in SC-Uracil media. Haploid cells coming from the unmodified parent cell show red pigment only when pCas9 plasmid was also present. In all cases, it was possible to PCR and sequence the barcode sequence from each haploid individual.

**Discussion**

Version control has been a pillar of software engineering and—withstanding that strain engineering is a very different discipline than software engineering—we believe that wider adoption of version control principles could substantially improve the quality of research that relies on modifying and engineering cell lines. We thus postulate that adoption of CellRepo will improve:

- Traceability: by physically linking a cell line chromosome to a commit id in the cloud, one can know the exact documentation for a strain. Besides technical information about the cell line, stored information also includes the intention behind genetic changes and allows proper allocation of credit (who created a particular commit in a cell line) for work done in the laboratory.
- Responsibility: because key cell lines can be tracked, branched, audited and ownership assigned both digitally and molecularly provenance, quality assurance and trust-worthiness are enhanced.
Reproducibility: it will be easier to reproduce experiments and avoid false leads because one will have a complete long-term change history of every modification to cell lines of interest. This change history includes the author of the change, laboratory of origin, the date of the change and written notes on the purpose and intention of each modification. Having a complete history of cell line modification also provides the ability to “revert” back to previous versions of a cell line, which is great for bug fixing in software engineering, and we believe will be useful in cell engineering too. As biologists, this would mean knowing exactly what someone else did at each commit-able stage in a project. Furthermore, this enables to base, with ease and confidence, a new cell line project on trustworthy pre-existing repositories and thus absorbing the history that came before it.

Collaboration: CellRepo improves collaboration. Version control systems allow complex software to be written by single individuals as well as by remote teams. Similarly, CellRepo accommodates both single or multi-scientist projects, maintaining a clear record of contributions. Furthermore, CellRepo does not force new workflows into laboratory users. Rather it is agnostic to the specific tools they already use and can accommodate uploads from any laboratory tools they may already be using. CellRepo also allows fine-tuning of a cell engineering project visibility by allowing repositories to be entirely private, shareable or public.

Transparency: our proposed version control system for cell engineering calls for more transparency in the process of making science. As we argued, research ought to be transparent and transparency benefits internal teamwork and enhances public trust in science. CellRepo empowers the sharing of cell line repositories in the same way that version control systems such as GitHub, Bitbucket or Gitlab host and promote open source projects. Like with open source projects, each snapshot of a cell repository shows the “good, the bad, and the ugly” of each stage in the development of a cell line. Through transparency, “bugs in the bugs” would be more readily discovered and corrected. Furthermore, as recently argued, there are two growing trends in science. One seeks to make science more open and the other more reproducible, but the adherents of these two trends do not always work concurrently toward openness and reproducibility. We believe our paper is a step toward bridging these two camps.

Economics: in software engineering, it is possible to develop software without using any version control. However, doing so subjects project owners to data and source code loss risk, loss of project history and the inability to collaborate in real-time. No professional software engineering team should or would accept those risks. Thus, we expect that as CellRepo become the norm in life sciences, important economic benefits will become more tangible.

We have adapted recombineering protocols to barcoding for version control in four bacterial species and two fungal species thus, in principle, one can create a truly universal tracking system for all lab-made cell lines. The barcodes introduced into key cell engineering milestones are the commit ids from the version control system bio-orthogonally mapped to DNA. We note that these barcodes fulfill a different role than whole-genome sequencing of a milestone, and hence cannot be solely replaced by it. For example, two different cell engineering projects might start from the same strain (hence having the same genome sequence) but require that they be distinguished from each other: different teams and laboratories, different goals and objectives for the project, different material transfer agreements or IP regimes, etc. Barcodes, watermarks and similar digital signatures embedded in the genome can be used to implement more sophisticated “digital rights management” that genome data by itself cannot. Moreover, although whole-genome sequencing is becoming cheaper, it is still far more expensive and complex process than sequencing a relatively short barcode as we propose here. Altogether, our work demonstrates that barcoding technology can be applied to many industrially and academically relevant microbial species; the barcode sequences are stable under laboratory conditions, they do not affect the growth of the barcoded strains and they can be used as a backtrack signal during GMO dispersion experiments. In the future, other kingdoms of life will also be added to CellRepo and tested in similar ways. Indeed, mammalian cells can be added straight away to CellRepo without a barcode or they could be barcoded via CRISPR/CAS methods or using lentiviral routes.

Importantly, we believe that a more traceable, reproducible and transparent development of engineered cell lines will contribute to improved public attitudes to the discipline. Indeed, public attitudes toward science in general, and in particular newly emerging technologies such as engineering biology, have been the focus of study over the last few decades. Some of the earlier studies were based on the so-called deficit model that assumes that the publics’ support of science and novel technologies is built on their knowledge of science. Although this model is still used by some scientists, surveys and polls have shown that more knowledge of science (or technology) does not necessarily lead to more public support. It was also shown that deference to science had a large impact on perceptions of, e.g., nanotechnology and that public trust in the claims made by experts or by scientific institutions is an important factor in understanding the publics’ attitudes to science. Similarly, although general support was found for the then-emerging field of synthetic biology, it was tempered by fears over misuse, health and environmental impacts, control and governance. Taken together these findings suggest that the publics’ concerns about engineering biology rest more with the method and processes underpinning the research rather than in the better understanding of one or another specific technical advance. That is, transparency and accountability of the research are the main concern in the publics’ attitudes to engineering biology.

Moreover, the potential for public mistrust in innovative science has been echoed in the science community’s crisis of conscience about the integrity of science itself. Similar concerns have been raised by other studies, UK Research and Innovation/Research England’s survey on research integrity confirmed that open and transparent research is regarded as central to rigor, reproducibility, and public trust. Commentators on the “crisis” broadly agree that the research community is overwhelmingly motivated by these values but that other factors within the research culture can make it difficult to uphold these principles (e.g., see). In response, there have been moves to promote measures that enable transparency, reproducibility and openness. For example, the “FAIR Principles”, namely findable, accessible, interoperable and re-usable, have been widely adopted while the UKRI Research Concordat on Open Research Data has been adopted by multiple stakeholders within the UK research community, with similar schemes elsewhere. The issues just highlighted also affect engineering biology and life sciences more generally.

Engineering biology, and in particular, strain engineering is hard, but the community working on cell engineering makes it harder by not adequately documenting, tracking and sharing the
process of genetic engineering. Baker describes the important contribution quality assurance processes can make to research integrity by enabling reproducibility and avoiding the opportunity for cherry-picking of results and data massaging. With this spirit in mind, in this paper, we have introduced a version control system for cell engineering. Versioning biological cells will lead to more trustworthy cell engineering.

Methods
Repository access. CellRepo can be accessed from https://cellrepo.ico2s.org

Creating or searching for a CellRepo repository. Repositories stored in CellRepo contain the entire digital footprint of a strain engineering process, which is linked to the cells in question via their genome stored barcode. Each repository represents an experiment/project on a cell line. To create a new repository, the user must navigate to the "Repositories" page using the navigation bar at the top of the page or the card on the homepage. Click the "Add" button to create a new repository. The repository can now be named and described in the form and the visibility of the project can be selected. The species of the repository can be chosen from a list of pre-selected species; if the desired species is not available in any of the linked databases then the user can request the addition of the new species. Finally, the repository can be created by clicking the "Create" button.

The user can also choose to start their experiments from a "branch" of another repository. To do this, they can search for project keywords or, importantly, barcode sequences using the "Search" functionality and create a new branch or commit from that point onwards.

Once the repository is created, the user can now start building their project using commits (see Fig. 4). Each commit adds the latest changes to the repository. It contains information about the changes to the cell line, who did it, references, barcodes, documents etc. Commits are a representation of a cell line at an exact moment in time. To create new commits, select "New Commit" from the Actions menu to add a commit to the repository. Fill in the commit form with information about your change. Add any files you want to upload that supplement the commit. At this step, the user can choose to add a DNA barcode by ticking a box. After synthesis, the DNA barcode can now be inserted into the strain’s genome by the protocols described in the Supplementary Information.

After more experiments are carried out on the strain, the repo can be updated, more information and documentation added, and the barcode sequence can be updated in the genome. These steps can be repeated as many times as necessary to document the history of the strain and the project.

Barcoding site selection. *E. coli* and *B. subtilis* have known lists of essential genes. Using these lists, it was possible to create a simple python script to get possible candidates of essential gene pairs. The script used as input the GFF3 annotation file of the strain and the list of the essential genes. Both files had to be curated to obtain a uniform gene name nomenclature. As output, the algorithm gives back a pair of essential genes next to each other, the orientation of both genes and the DNA sequence that separates them. Then, databases and prediction tools were used to check for the presence of regulatory elements in the intergenic sequences that did not appear in the annotation files. Once a good candidate pair was obtained, the target region was aligned against the most common laboratory strains of the species to check for the presence of the possible barcoding region in them.

No list of core essential genes for *P. putida* strains was found in the literature except for conditional essential genes in some conditions and essential genes in the related species *Pseudomonas aeruginosa*. For this reason, well-known generally conserved essential genes were taken into consideration to choose one possible insertion locus for the barcode. *glmS* gene was chosen as a good candidate as it is a broad-host conserved gene in many species and it has a long enough intergenic region for the insertion of Ll.LtrB intron carrying barcodes. The procedure to select the exact insertion site inside the intergenic region downstream of *glmS* was adapted from. In general, the PP5408-*glmS* intergenic region was surveyed for good Ll.LtrB intron insertion sites in the Clostron.com website. The sequences needed for tn7 insertion were avoided as we did not want to hinder the possibility of using this insertion method before or after the barcoding procedure. From the retrieved list of insertion loci, one was chosen from previous data verifying the correct insertion of Ll.LtrB in this site.
Similarly, to *P. putida*, no essential gene information was found for *S. albidoflavus* (previously known as *Streptomyces albus*). However, in this case, we wanted to follow a "cost-less" approach to showcase the flexibility of the proposed system for new candidate species to be barcoded. A close relative, *S. coelicolor*, is the model species for the study of the Streptomyces genus. For this species, there is a genome-scale metabolic model with gene essentiality data available that can be applied to other *Streptomyces* species. Using this model, just reactions catalyzed by a known *S. coelicolor* gene, essential for all conditions tested and with no isoenzymes, were considered. Using this data, it was possible to create a list of putative essential genes (Supplementary Table 1) for *S. albidoflavus* by aligning each of the *S. coelicolor* A3(2) essential genes to *S. albidoflavus* i1074 database. Feeding this list to the script described found no good candidate pair. The number of essential genes next to each other was too low and the few candidate pairs found had problematic intergenic regions. A simpler approach was followed.

Using the list of putative essential genes for i1074, the whole genome was analyzed looking for clusters of nearby essential genes. In this case, the resulting candidate pair genes were not next to each other but separated by one or more non-essential genes.

To check if the putative essential gene pair was conserved among different *Streptomyces* species, the protein sequences of the chosen candidate genes were aligned against the *Streptomyces* protein database using BLAST to check if the genes were conserved among different species (Supplementary Fig. 1). The results suggest that the two selected genes are conserved and are good candidates for essentiality.

*S. cerevisiae* is well known among the synthetic biology community and there is plenty of well-curated information about it. The user of CellRepo could choose to insert the barcode sequence into an already known and curated insertion site. These sites are used for example in microbial cell factories to experiment with essential heterologous genes. By using this type of site, the possibility to barcode a strain, while the user’s desired edition occurs is shown feasible. It was decided to go for an already known insertion site flanked by essential genes used in microbial cell factories experiments\(^2\). Also, this site has previously been used to test CRISPR plasmids in *S. coelicolor*\(^3\).

*K. phafii* (previously known as *P. pastoris*) is known for its ability to produce high amounts of recombinant protein. The alcohol oxidase *AOX1* promoter insertion site is commonly used because of its tight regulation and strength\(^4\). For these reasons, the *AOX1* promoter site was selected as the insertion site in *K. phafii* without considering closeness to essential genes.

**Barcoding protocols.** Different barcoding methods were designed for each species. For this study, all the selection markers were removed from the final strains, except for *K. phafii*.

For *E. coli*, *B. subtilis*, *S. albidoflavus*, *S. cerevisiae* and *K. phafii* vectors containing a restriction site to allow the easy cloning of the barcode sequence by Hi-Fi assembly (NEB) using restriction-linearized vectors were used. The barcode DNA sequences were synthesized as dsDNA fragments (Integrated DNA Technologies) and inserted into the plasmids. The barcoding vectors of *P. putida* were based on the procedure described\(^5\) adapted to this microorganism\(^6\). Please see Supplementary Table 2 and Supplementary Fig. 2 for a detailed description of the vectors used in this study.

To simplify experiments for this paper, just one DNA sequence was used to barcode each species except for *S. cerevisiae* which each of the two barcoding procedures used a different barcode (Supplementary Table 3). In a real-life scenario, the usage of CellRepo produces different DNA sequence barcodes for each commit (to avoid clashes and ambiguity).

**Supplementary Information** contains a detailed description of the protocols used to barcode each species and the different growth media used in our studies.

**Growth curves.** For these experiments, all selection markers inserted in the strains were removed (except for *K. phafii*). All the growth curve experiments were carried out in a CLARIOstar® Plus (BMG Labtech) plate reader using a polystyrene sterile plate, at 300 rpm, using three biological replicates per strain. *E. coli* and *B. subtilis* were grown in LB medium at 37°C measuring the absorbance at 600 nm. *P. putida* was grown in LB, at 30°C. *S. cerevisiae* and *K. phafii* cells were grown at 30°C in YPD medium in 24-well plates. *S. albidoflavus* plate reader experiment was carried out in TS-agar as previously described\(^7\) for *S. coelicolor*.

**Whole-genome sequencing.** The genomic DNA was extracted using GenElute® Bacterial Genomic DNA Kit Protocol (Sigma) (*E. coli*, *B. subtilis*, *P. putida* and *S. albidoflavus*) and YeastStar Genomic DNA Kit (Zymo Research) (*S. cerevisiae* and *K. phafii*).

NGS library was prepared using NEB Next® Ultra® DNA Library Prep Kit (Cat No. E7370L). Whole-genome sequencing was performed on an Illumina NovaSeq 6000 platform at Novogene (Beijing, China).

The reads were aligned against the reference genome of each species (Supplementary Table 3) using Geneious Prime 2019.2.3 (https://www.geneious.com). First, reads were trimmed using BBduk (Adapter/Quality Trimming Version 38.37 by Brian Bushnell) and then duplicates were removed using Dedupe (Duplicate Read Remover 38.37 by Brian Bushnell) with default settings in Geneious. The reads were then mapped against the reference genomes using the following settings: Mapper “Genious”, Sensitivity “Medium/Low/Fast” and selecting the “Find structural variants of any size” option. To annotate SNPs, Geneious integrated algorithm with default settings and a “Minimum variant frequency” value of 0.5 was used. All the mutations also found in the wild-type strain genome were discarded from the analysis following the next steps:

1. Use the comparison tool in Geneious to remove all the SNPs present in the wild-type strains.
2. Manually curate the rest of SNPs, focusing especially on low coverage and repetitive regions (frequent in both *S. cerevisiae* and *K. phafii*).

SNPs flagged in barcoded strains are not in wild-type strains and vice versa. This is because some detected variants qualify as such in one strain but not in the other due to coverage, quality, etc.

**Barcode survival.** The stability of the barcode sequence was tested by growing each species for 10 days under different growth conditions.

**Condition 1: 10 ml of the growth media (LB for *E. coli*, *B. subtilis* and *P. putida*; TSB for *S. albidoflavus*; YPD for *S. cerevisiae* and *K. phafii*) were inoculated and grown overnight at 200 rpm. Each morning, during the following 10 days, 100 ml of the culture were re-inoculated in 10 ml of fresh media.**

**Condition 2: 50 ml of the growth media were inoculated and grown at 200 rpm for 10 days.**

**Condition 3: using a BioXplorer 400 (HEL, London) bioreactor, the cells were grown in a 75 ml of growth media with impeller agitation (400 rpm) and filtered air supply (100 ml/min). Cells were grown overnight. After the first night, cells were grown continuously for 10 days at a dilution rate (D) of 0.024 h⁻¹ (minimum possible setting of the system). Antifoam 204 (Sigma A6426) was added to the liquid media before autoclaving at 0.01%.**

**Condition 4: using the same bioreactor system described in the previous condition cells were grown overnight. After the first night, cells were grown continuously for 10 days at a dilution rate (D) of: 0.3 h⁻¹ for *E. coli*, *B. subtilis* and *P. putida*; 0.2 h⁻¹ for *S. albidoflavus*, *S. cerevisiae* and *K. phafii*. The dilution rates were inferred from commonly used values for continuous culture\(^8\) and previously described growth curves.**

**Condition 5: three colony replicates of the barcoded strains were restreaked on solid media for ten passes.**

For conditions 1–4, samples were taken periodically and plated on solid media to ensure no contamination had occurred. On the last day of the experiments, a sample was taken and spread on agar plates of the same growth media. Single colonies were restreaked. For all conditions, the barcode region was amplified by PCR after genomic DNA extraction and sent to Sanger sequencing (Eurofins Genomics). The sequencing results were aligned against the reference barcode sequence for each species.

**The genomic DNA of the microbial population of conditions 1–4 were isolated as previously described. Also, as a control experiment that did not go through the 10-day culture period, the gDNA of the glycerol stock used to start the long-term experiments was extracted as well. The barcode region was amplified by PCR and the amplicon was used for NGS analysis. DNA library preparations, sequencing reactions, and initial bioinformatics analysis were conducted at GENEWIZ, Inc. (South Plainfield, NJ, USA). DNA amplicons with partial adapters were indexed and enriched by limited cycle PCR. The DNA library was validated using TapeStation (Agilent Technologies, Palo Alto, CA, USA), and was quantified using Qubit 2.0 Fluorometer and real-time PCR (Applied Biosystems, Carlsbad, CA, USA). The pooled DNA libraries were loaded on the Illumina instrument according to the manufacturer’s instructions. The samples were sequenced using a 2 × 250 paired-end configuration. Image analysis and base calling were conducted by the Illumina Control Software (HCS) on the Illumina instrument.**

**Raw Fastq data were first trimmed to remove low-quality data using sickle (https://github.com/najoshi/sickle). PANDAseq (https://github.com/neufeld/pandaseq)\(^7\) was then used to merge read1 and read2 of each sample. The merged reads of each sample were mapped to the target reference sequence using BWA (http://bio-bwa.sourceforge.net)\(^7\). Then variants were detected using GENEWIZ’s in-house script. The primers used for NGS amplicon analysis can be found in Supplementary Table 4.**

**Yeast gene drive.** Special contingency and sterility measures were taken to perform the gene drive experiments. All the experiments were carried out in a Class 2 safety cabinet. All the surfaces were UV-light and chemically sterilized. All the agar plates were sealed using Parafilm. pGD-AD2 was assembled containing homologous regions to AD2 gene, URA3 marker (from *B.5-Ura3*), a sgRNA targeting AD2 and a barcode sequence (both chemically synthesized as gblocks). A modified version of the protocol detailed in Reference 75 was followed. Briefly, the PCR amplification product of the previous cassette was transformed into BY4741 cells. pCB23212 (Cas9) plasmid
Table 1 List of CellRepo-hosted public supplementary data.

| Species     | Link to CellRepo                                                                 | QR code |
|-------------|----------------------------------------------------------------------------------|---------|
| E. coli     | https://cellrepo.ico2s.org/repositories/59?branch_id=826&locale=en              |         |
| B. subtilis | https://cellrepo.ico2s.org/repositories/60?branch_id=85&locale=en               |         |
| P. putida   | https://cellrepo.ico2s.org/repositories/61?branch_id=89&locale=en               |         |
| S. albidoflavus | https://cellrepo.ico2s.org/repositories/62?branch_id=91&locale=en         |         |
| S. cerevisiae | https://cellrepo.ico2s.org/repositories/65?branch_id=96&locale=en           |         |
| K. phaffii  | https://cellrepo.ico2s.org/repositories/64?branch_id=94&locale=en             |         |

This table provides, for each of the species used in this paper, a URL and a QR code that point to the public repository containing the data associated with each species-specific barcoding protocol.

was transformed afterwards. BY4741-GeneDrive-pCB8312 cells were mated with BY4742-pXP622 (used just to select diploids) and plated in SC-Leucine containing G418 (200 μg/mL). Diploid cells were then grown in GNA solid media overnight. Cells were transferred to SPOR plates and sporulated following the protocol described in[3]. Briefly, SPOR plates were incubated at room temperature overnight and 30 °C for 5 days. When tetrads were observed, some cells were scraped from the plate and cell wall digested using Zymolyase solution after incubation at 37 °C for 20 min. Tetrads were dissected using SporePlay (+ Sanger Instruments). Single spores were grown on YPD plates until colony formation. Cells were resuspended in water and 5 μL were transferred to GNA and SC-Uracil plates.

pXP622 was a gift from Nancy Dahlva & Suzanne Sandmeyer (Addgene plasmid # 26849). BS-Ura3Kl was a gift from Zhiping Xie (Addgene plasmid # 894&locale=en)

K. phaffii https://cellrepo.ico2s.org/repositories/64?branch_id=94&locale=en

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

Data availability

The authors declare that data supporting the findings of this study are available within the paper and its Supplementary Information files. DNA sequencing data are available at NCBI under accession PRJNA797888. In addition, other protocols are publicly available at https://cellrepo.ico2s.org/got as well as at the repositories listed in Table 1.

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References

1. Jinek, M. et al. A programmable dual-RNA-guided DNA endonuclease in adaptive bacterial immunity. Science 337, 816–821 (2012).
2. Mehlhenbacher, R. D., Kolbl, R., Lay, A. & Dionne, J. A. Nanomaterials for in vivo imaging of mechanical forces and electrical fields. Nat. Rev. Mater. 3, 17080 (2017).
3. Farhadi, A., Sigmund, F., Westmeyer, G. G. & Shapiro, M. G. Genetically encodable materials for non-invasive biological imaging. Nat. Mater. 20, 585–592 (2021).
4. Gilbert, C. et al. Living materials with programmable functionalities grown from engineered microbial co-cultures. Nat. Mater. 20, 691–700 (2021).
5. de Lorenzo, V. et al. The power of synthetic biology for bioproduction, remediation and pollution control. EMBO Rep. 19, e56588 (2018).
6. Yin, S. S. et al. Robust direct digital-to-biological data storage in living cells. Nat. Chem. Biol. 17, 246–253 (2021).
7. Morrison, C. & Lähteenmäki, R. Public biotech in 2017—the numbers. Nat. Publ. Gr. 36, 576–584 (2018).
8. Vogt, C. A. Synthetic biology 2020–2030: six commercially-available products that are changing our world. Nat. Commun. 11, 6379 (2020).
9. Zhou, P. et al. A pneumonia outbreak associated with a new coronavirus of probable bat origin. Nature 579, 270–273 (2020).
10. Anzalone, A. V. et al. Search-and-replace genome editing without double-strand breaks or donor DNA. Nature 576, 149–157 (2019).
11. Blakes, J. et al. Heuristic for maximizing DNA reuse in synthetic DNA library assembly. ACS Synth. Biol. 3, 529–542 (2014).
12. Yehezkel, T. Ben et al. Synthesis and cell-free cloning of DNA libraries using programmable microfluidics. Nucleic Acids Res. 44, e35 (2016).
13. Reardon, S. CRISPR gene-editing creates waves of exotic model organisms. Nature 568, 441–442 (2019).
14. de Lorenzo, V., Krasnogor, N. & Schmidt, M. For the sake of the bioeconomy: define what a synthetic biology chassis is! N. Biotechnol. 60, 44–51 (2021).
15. Identity crisis. Nature 457, 935–936 (2009).
16. American Type Culture Collection Standards Development Organization & Workgroup ASN-0002. Cell line misidentification: the beginning of the end. Nat. Rev. Cancer 10, 441–448 (2010).
17. Masters, J. R. End the scandal of false cell lines. Nature 492, 186 (2012).
18. Challenges in irreproducible research. Nature collection. www.nature.com/collections/prbfkwmwvz/ (2018).
19. Mehra, M. R., Desai, S. S., Ruschitzka, F. & Patel, A. N. RETRACTED: hydroxychloroquine or chloroquine with or without a macrolide for treatment of COVID-19: a multinational registry analysis. Lancet 1–10 https://doi.org/10.1016/S0140-6736(20)31180-6 (2021).
20. Tellechea-Luzzardo, J. et al. Linking engineered cells to their digital twins: a version control system for strain engineering. ACS Synth. Biol. 9, 536–545 (2020).
21. Committee, E. S. et al. Evaluation of existing guidelines for their adequacy for the microbial characterisation and environmental risk assessment of microorganisms obtained through synthetic biology. EFSA J 18, e06263 (2020).
22. Schmidt, M. & de Lorenzo, V. Synthetic bugs on the loose: containment options for deeply engineered (micro)organisms. Curr. Opin. Biotechnol. 38, 90–96 (2019).
23. Beeckman, D. S. A. & Rüdelsheim, P. Biosafety and biosecurity in containment: a regulatory overview. Front. Bioeng. Biotechnol. 8, 650 (2020).
24. Baig, H. et al. Synthetic biology open language (SBOL) version 3.0.0. J. Integr. Bioinform. 17, 20200017 (2020).
25. Stovicke, V., Holkenbrink, C. & Borodina, I. CRISPR/Cas system for yeast genome engineering: advances and applications. FEMS Yeast Res. 17, fox030 (2017).
26. Tong, Y., Charusanti, P., Zhang, L., Wang, T. & Lee, S. Y. CRISPR-Cas9 based engineering of actinomycetal genomes. ACS Synth. Biol. 4, 1020–1029 (2015).
27. Wijsman, M. et al. A toolkit for rapid CRISPR-SpCas9 assisted construction of hexose-transport-deficient Saccharomyces cerevisiae strains. FEMS Yeast Res. 19, fox107 (2019).
28. Ryan, F. J., Nakada, D. & Schneider, M. J. Is DNA replication a necessary condition for spontaneous mutation? Z. Vererbungsl. 91, 38–41 (1961).
29. Ryan, F. J., Okada, T. & Nagata, T. Spontaneous mutation in spheroplasts of Escherichia coli. J. Gen. Microbiol. 30, 153–158 (1963).
30. Loewe, L., Textor, V. & Scherer, S. High deleterious genomic mutation rate in stationary phase of Escherichia coli. Science 302, 1558–1560 (2003).
31. Bull, H. J., Mckenzie, G. J., Hastings, P. J. & Rosenberg, S. M. Evidence that stationary-phase hypermutation in the Escherichia coli chromosone is promoted by recombination. Genetics 154, 1427–1437 (2000).
32. Sun, H.-M. & Yasbin, R. E. Adaptive, or stationary-phase, mutagenesis, a component of bacterial differentiation in Bacillus subtilis. J. Bacteriol. 184, 5641–5653 (2002).
33. Kasak, L., Hörak, R. & Kivisaar, M. Promoter-creating mutations in Pseudomonas putida: a model system for the study of mutation in starving bacteria. Proc. Natl. Acad. Sci. USA. 94, 3134–3139 (1997).

34. Steele, D. F. & Jinks-Robertson, S. An examination of adaptive reversion in Saccharomyces cerevisiae. Genetics 132, 9–21 (1992).

35. Achilli, A. et al. The exceptionally high rate of spontaneous mutations in the polymerase delta proofreading exonuclease-deficient Saccharomyces cerevisiae strain starved for adenine. BMC Genet. 5, 34 (2004).

36. Noble, C. et al. Daisy-chain gene drives for the alteration of local populations. Proc. Natl. Acad. Sci. USA. 116, 8275–8282 (2019).

37. Windbichler, N. et al. A synthetic homing endonuclease-based gene drive system in the human malaria mosquito. Nature 473, 212–215 (2011).

38. Murphy, M. C. et al. Open science, communal culture, and women’s participation in the movement to improve science. Proc. Natl. Acad. Sci. USA. 117, 24154–2412164 (2020).

39. Akin, H. et al. Mapping the landscape of public attitudes on synthetic biology. Bioscience 67, 290–300 (2017).

40. Sturpis, G. & Allum, N. Science in society: re-evaluating the deficit model of public attitudes. Public Underst. Sci. 13, 55–74 (2004).

41. Ho, S. S., Scheufele, D. A. & Corley, E. A. Making sense of policy choices: understanding the roles of value predispositions, mass media, and cognitive processing in public attitudes toward nanotechnology. J. Nanopart. Res. 12, 2703–2715 (2010).

42. Jasanoff, S. The ‘science wars’ and American politics. In Between Understanding and Trust (eds Dierkes M. & Grote von C.) 14 (Routledge, 2000).

43. Yearley, S. What does science mean in the ‘public understanding of science’. (2000).

44. Bhattachary, D., Pascall Calitz, J. & Hunter, A. Synthetic biology dialogue. Science academies, Science and Trust, Executive summary and recommendations. Summit G7 (2019).

45. Ho, S. S., Scheufele, D. A. & Corley, E. A. Making sense of policy choices: understanding the roles of value predispositions, mass media, and cognitive processing in public attitudes toward nanotechnology. J. Nanopart. Res. 12, 2703–2715 (2010).

46. Jasanoff, S. The ‘science wars’ and American politics. In Between Understanding and Trust (eds Dierkes M. & Grote von C.) 14 (Routledge, 2000).

47. Baker, M. 1,500 scientists lift the lid on reproducibility. Nature 529, 290–300 (2015).

48. Gromova, M. et al. How quality control could save your science. BMC Biotechnol. 5, 34 (2004).

49. Nosek, B. A. et al. Promoting an open research culture. Proc. Natl Acad. Sci. USA. 115, 3922–3924 (2018).

50. What researchers think about the culture they work in. Nature 2016, 3134–3139 (1997).

51. The Concordat on Open Research Data. Nature 2017, 3134–3139 (1997).

52. Baker, M. How quality control could save your science. Nature 529, 456–458 (2016).

53. Baker, M. How quality control could save your science. Nature 529, 456–458 (2016).

54. Ballesta, T. et al. Construction of Escherichia coli K-12 in-frame, single-gene knockout mutants: the Keio collection. Mol. Syst. Biol. 2, 692–698 (2014).

55. Karp, P. D. et al. The BioCyc collection of microbial genomes and metabolic pathways. Brief. Bioinform. 20, 1085–1093 (2017).

56. Solovyev, V. & Salamov, A. Automatic annotation of microbial genomes and metagenomic sequences. In Metagenomics and its Applications in Agriculture. (ed. Robert W. Li), (2010).

57. Lesnik, E. A. et al. Prediction of rho-independent transcriptional terminators in Escherichia coli. Nucleic Acids Res. 29, 3583–3594 (2001).

58. Molina-Haenares, M. A. et al. Identification of conditionally essential genes for growth of Pseudomonas putida KT2440 on minimal medium through the screening of a genome-wide mutant library. Environ. Microbiol. 12, 1468–1485 (2010).

59. Liberati, N. T. et al. An ordered, nonredundant library of Pseudomonas aeruginosa strain PA14 transposon insertion mutants. Proc. Natl. Acad. Sci. USA. 103, 28333–28339 (2006).

60. Poulsen, B. E. et al. Defining the core essential genome of Pseudomonas aeruginosa. Proc. Natl. Acad. Sci. USA. 116, 10072–10073 (2019).

61. Lee, S. A. et al. General and condition-specific essential functions of Pseudomonas aeruginosa. Proc. Natl Acad. Sci. USA. 112, 5189–5194 (2015).

62. Velázquez, E., Lorenzo, Vde & Al-Ramahi, Y. Recombination-independent genome editing through CRISPR/Cas9-enhanced TargeTron delivery. ACS Synth. Biol. 8, 2186–2193 (2019).

63. Velázquez, E., Al-Ramahi, Y., Tellechea, J., Krasnogor, N. & de Lorenzo, V. Targeted-assisted delivery of exogenous DNA sequences into Pseudomonas putida through CRISPR-aided counterselection. ACS Synth. Biol. 10, 2552–2565 (2021).

64. Borodina, I., Krabben, P. & Nielsen, J. Genome-scale analysis of Streptomyces coelicolor A3(2) metabolism. Genome Res. 3, 820–829 (2005).

65. Dalggaard Mikkelsen, M. et al. Microbial production of indolylglycosaminolysine through engineering of a multi-gene pathway in a versatile yeast expression platform. Metab. Eng. 14, 104–111 (2012).

66. Jessop-Fabre, M. M. et al. EasyClone-MarkerFree: a vector toolkit for markerless integration of genes into Saccharomyces cerevisiae via CRISPR-Cas9. Biotechnol. J. 11, 1110–1117 (2016).

67. Ahmad, M., Hirz, M., Pichler, H. & Schwab, H. Protein expression in Pichia pastoris: recent achievements and perspectives for heterologous protein production. Appl. Microbiol. Biotechnol. 98, 5301–5317 (2014).

68. Santos-Benet, F. Genome sequencing analysis of Streptomyces coelicolor mutants that overcome the phosphate-dependent vancomycin lethal effect. BMC Genomics 19, 457 (2018).

69. Peebo, K. & Neubauer, P. Application of continuous culture methods to recombinant protein production in microorganisms. Microorganisms 6, 56 (2018).

70. Masella, A. P., Bartram, A. K., Truszkowski, J. M., Brown, D. G. & Neufeld, J. D. PANDAseq: PAired-EDN assembler for illumina sequences. BMC Bioinformatics 13, 31 (2012).

71. Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows-Wheeler transformer. Bioinformatics 25, 1754–1760 (2009).

72. Li, D. et al. A fluorescent tool set for yeast Atg proteins. Autophagy 11, 954–960 (2015).

73. D'Carlo, J. E., Chavez, A., Dietz, S. L., Esvelt, K. M. & Church, G. M. Safeguarding CRISPR-Cas9 gene drives in yeast. Nat. Biotechnol. 33, 1250–1255 (2015).

74. Treco, D. A. & Winston, F. Growth and manipulation of yeast. Curr. Protoc. Mol. Biol. Chapter 13, Unit 13.2 https://doi.org/10.1002/0471127277.mb130282 (2008).

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Conceptualization: N.K.; methodology: J.T.L., E.V., L.H., V.d.L., N.K.; investigation: J.T.L., E.V., L.H.; software: L.H., N.K.; writing—original draft: J.T.L., L.H., E.V., L.P., S.W., V.d.L., N.K.; supervision: N.K., V.d.L.; project administration: N.K.; funding: N.K., S.W., V.d.L.

Competing interests

The authors declare no competing interests.

Additional information

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