The genome of *Schmidtea mediterranea* and the evolution of core cellular mechanisms

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The planarian *Schmidtea mediterranea* is an important model for stem cell research and regeneration, but adequate genome resources for this species have been lacking. Here we report a highly contiguous genome assembly of *S. mediterranea*, using long-read sequencing and a de novo assembler (MARVEL) enhanced for low–complexity reads. The *S. mediterranea* genome is highly polymorphic and repetitive, and harbours a novel class of giant retroelements. Furthermore, the genome assembly lacks a number of highly conserved genes, including critical components of the mitotic spindle assembly checkpoint, but planarians maintain checkpoint function. Our genome assembly provides a key model system resource that will be useful for studying regeneration and the evolutionary plasticity of core cell biological mechanisms.

Rapid regeneration from tiny pieces of tissue makes planarians a prime model system for regeneration. Abundant adult pluripotent stem cells, termed neoblasts, power regeneration and the continuous turnover of all cell types1–3, and transplantation of a single neoblast can rescue a lethally irradiated animal4. Planarians therefore also constitute a prime model system for stem cell pluripotency and its evolutionary underpinnings5. The taxonomic clade Platyhelminthes (‘flatworms’) also includes parasitic lineages that have substantial effects on human health, such as blood flukes (Trematoda) and tape worms (Cestoda)6. Here, the phylogenetic position of planarians as free-living flatworms provides a reference point towards an understanding of the evolution of parasitism8.

Despite the modest genome sizes of planarians (mostly in the range of 1–2 gigabase pairs (Gb)), genome resources relating to these animals are limited. Although the model species *S. mediterranea* was sequenced by Sanger sequencing, even 11.6× coverage of around 600-bp Sanger reads yielded only a highly fragmented assembly (N50 19 kb)9. Recent high-coverage, short-read approaches yielded similarly fragmented assemblies10,11. The high A–T content (about 70%) represents one known assembly challenge. Furthermore, standard DNA isolation procedures perform poorly on planarians, which has so far precluded the application of long-read sequencing approaches or BAC-clone scaffolding.

We here report a highly contiguous PacBio SMRT long-read sequencing assembly of the *S. mediterranea* genome. Giant gypsy/Ty3 retroelements, abundant AT-rich microsatellites and inbreeding-resistant heterozygosity collectively provide an explanation for why previous short-read approaches were unsuccessful. We find a loss of gene synteny in the genome of *S. mediterranea* and other flatworms. In analysis of highly conserved genes, we find a loss of MAD1 and MAD2, suggesting a MAD1–MAD2-independent spindle assembly check point (SAC)13,14. Our *S. mediterranea* genome assembly provides a resource for probing the evolutionary plasticity of core cell biological mechanisms, as well as the genomic underpinnings of regeneration and the many other phenomena that planarians expose to experimental scrutiny.

**De novo long read assembly of the planarian genome**

In preparation for genome sequencing, we inbred the sexual strain of *S. mediterranea* (Fig. 1a) for more than 17 successive sib-mating generations in the hope of decreasing heterozygosity. We also developed a new DNA isolation protocol that meets the purity and high molecular weight requirements of PacBio long-read sequencing12 (Extended Data Fig. 1a–d, Supplementary Information S1, S2). We used MARVEL, a new long-read genome assembler developed for low complexity read data13 (Supplementary Information S3). An initial de novo MARVEL assembly of reads of more than 4 kb with approximately 60× genome coverage showed an improvement over the PacBio assembly tool (Canu15) and substantial improvements over existing *S. mediterranea* assemblies based on short read sequencing (Extended Data Table 1).

To assess the quality of this genome assembly, we back-mapped a transcriptome of the sequenced strain (Supplementary Information S6) and found that more than 99% of transcripts were mapped, thus confirming that the assembly was both near-complete and accurate (Supplementary Information S7, Extended Data Fig. 1f, g). To assess the contiguous of the global assembly, we analysed structural conflicts between the MARVEL assembly and Chicago/HiRise scaffolding. Out of 51 such events across the 782.1 Mb of assembled genome sequence, only two represented unambiguous MARVEL assembly mistakes (Fig. 1b, Supplementary Information S4.3). Furthermore, high-stringency back-mapping of high-confidence cDNA sequences (Supplementary Information S7.3) confirmed assembly contiguity below the approximately 1-kb resolution limit of the Chicago/HiRise method, with small-scale sequence duplications near assembly gaps as only minor inconsistencies (Extended Data Fig. 2).

Our *S. mediterranea* genome assembly represents a major improvement over existing *S. mediterranea* assemblies10 (Fig. 1c) and, to our

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knowledge, is the first long-range contiguous assembly of the genome of a non-parasitic flatworm species. A UCSC genome browser instance with supplementary quality control, annotation and experimental data tracks (Supplementary Information S8) is available at PlanMine18 (http://planmine.mpi-cbg.de). All analyses in this manuscript refer to the assembly release version dd_Smed_g4. The current source code of the MARVEL assembler is available at https://github.com/schloi/MARVEL. The execution scripts used for S. mediterranea can be found in the smed subfolder of the examples folder.

**Assembly challenges in the S. mediterranea genome**

To understand why the S. mediterranea genome was recalcitrant to earlier short-read assembly, we first analysed its repeat content (Supplementary Information S9). The genome has a repetitive fraction of 61.7% (Fig. 2a), substantially exceeding the 38% or 46% repeat content of the mouse or human genomes, respectively.19 We detected more than 7,000 insertions of 11 distinct families of long terminal repeat (LTR) retroelements (Fig. 2b, Extended Data Fig. 3a, Supplementary Information S10). These do not cluster with known Metaviridae (Fig. 2b), suggesting that they represent either extremely divergent or so far undescribed retroelement families. Three LTR families were more than 30 kb long—an exceptional size that is more than three times longer than the 5–10 kb typically observed in vertebrates (Fig. 2c, Extended Data Fig. 3b). The only known similar-sized LTRs are the plant-specific Ogre elements20, which is why we refer to the giant S. mediterranea repeat families as Burro (big, unknown rivaling ogre; Supplementary Information S10.3). Burro elements are pervasively transcribed (Extended Data Fig. 3c, d, Supplementary Information S10.4), yet their high degree of intra-family sequence divergence suggests a relatively ancient invasion (Supplementary Table 1, Supplementary Information S10.5, Extended Data Fig. 3e). Burro-1, the most abundant giant retroelement with 130 fully assembled copies, is highly overrepresented at contig ends, and 50% of all current scaffolds terminate in a Burro-1 element (Fig. 2d, Supplementary Information S10.6). Therefore, these abundant, over 30-kb repeat elements still limit the size of the current assembly. In addition, abundant AT-rich microsatellite regions disrupt the alignment of spanning reads and thus also reduce contig contiguity (Extended Data Fig. 4, Supplementary Information S11). Finally, the S. mediterranea assembly graphs showed substantial structural heterogeneity (Supplementary Information S12) in the form of bubbles (transient divergences in sequencing read alignments) and spurts (divergences without re-connection), which were largely absent from a comparable genome assembly (Drosophila melanogaster using PacBio sequencing and MARVEL assembly; Fig. 2e, Supplementary Information S12.1) or assemblies of 17 other species (Supplementary Table 2). Heterozygous mobile element insertions and microsatellite tracts were prominent causes of assembly divergences (Fig. 2f, Extended Data Fig. 4d, Supplementary Information S12.3). The persistence of substantial genomic heterozygosity in spite of more than 17 successive sib-mating generations confirms that meiotic recombination is inefficient in S. mediterranea21.

Overall, the combination of giant repeat elements, low-complexity regions and inbreeding-resistant heterozygosity provides an explanation for why previous short-read sequencing assemblies of S. mediterranea have proven so challenging. The long-range contiguity that we achieved in the S. mediterranea genome assembly, and similarly substantial improvements in the PacBio genome assembly of the flatworm Macrostomum lignano22 (Supplementary Table 2), further emphasize the improvements that a combination of long-read sequencing with the MARVEL assembler offers in the assembly of challenging genomes.

**Comparative analysis of the planarian gene complement**

We next annotated the S. mediterranea gene complement, relying on our planarian transcriptome resources18 (Supplementary Information S13). Our analysis showed a high divergence of S. mediterranea gene sequences (Supplementary Information S14), en par with Caenorhabditis elegans (Fig. 3a). By contrast, the low degree of sequence substitutions between the sexual and asexual S. mediterranea strains (Fig. 3a) and nearly identical mapping statistics of the two transcriptomes to the genome (Supplementary Information S7.1, Extended Data Fig. 1f) establish the utility of our assembly for both strains.

To evaluate the S. mediterranea genome structure, we performed whole-genome alignments (Supplementary Information S15) with the available parasitic flatworm genomes2 and a draft genome of the platyhelminth Macrostomum lignano22 (Fig. 3b). The highest alignment similarity was found between S. mediterranea and the parasitic flatworm Schistosoma mansoni, which is consistent with the platyhelminth phylogeny23. However, alignments were mostly limited to individual exons of specific genes, irrespective of the quality of the various assemblies (Extended Data Fig. 5a, b). In general, flatworm genome comparisons resulted in alignment chains that were much shorter and lower scoring than those obtained from comparisons across the tetrapod (human–frog) or vertebrate (human–zebrafish) clades (Fig. 3b). Together with more than 1,000 likely planarian-specific protein coding genes (Supplementary Information S16, Supplementary Table 5, Extended Data Fig. 6a–g), our data show a high degree of genome divergence between S. mediterranea and other flatworms.

We therefore next investigated gene loss in planarians. Our analysis deliberately focused on highly conserved genes, such that the absence of sequence similarity alone provides a strong indication of
loss (Supplementary Information S17). We identified 452 highly conserved genes that were lost in both S. mediterranea and other planarians (Fig. 3c), which compares to 284 and 757 such losses in D. melanogaster and C. elegans, respectively (Extended Data Fig. 5c). Gene loss in planarians is therefore broadly in the same range as in established invertebrate model organisms. However, the lost genes included 124 homologs that were lost in both S. mediterranea and other planarians (Supplementary Information S17). We identified 452 highly conserved genes that were lost in both S. mediterranea and other planarians (Supplementary Information S17). We identified 452 highly conserved genes that were lost in both S. mediterranea and other planarians (Supplementary Information S17). We identified 452 highly conserved genes that were lost in both S. mediterranea and other planarians (Supplementary Information S17). We identified 452 highly conserved genes that were lost in both S. mediterranea and other planarians (Supplementary Information S17).

Figure 2 | S. mediterranea assembly challenges. a. Repeat content of the assembly. b, LTR family phylogeny. Known LTR families are shown in colour, S. mediterranea LTR families in black. Red arcs delimit clusters for consensus calculation. Scale bar: 0.2 substitutions per site. c, Domain annotation of the 11 S. mediterranea LTR families (SLFs). d, Enrichment analysis of indicated repeat elements within the terminal 1,000 bp of all scaffolds (n = 962). Expected values represent mean repeat frequency with 95% bootstrap confidence interval (n = 1,000). e, Graphical representation of representative S. mediterranea (left, ~1.6 Mb) and D. melanogaster (right, ~1.7 Mb) MARVEL PacBio assembly graph segments. Thick lines, despite maintained haem biosynthesis capacity, is similarly unusual for a free-living eukaryote (both are lost in C. elegans). Remarkably, the above and multiple other genes were missing not only in planarians, but also in the parasite genomes and the transcriptome of the macrostomid M. lignano (Fig. 3c). Given their broad conservation in the lophotrochozoan sister clade, the broad absence of these genes in flatworms is likely to represent an ancestral loss. This complicates, for example, the interpretation of FASN loss in the parasitic lineages as a specific adaptation to parasitism. Conversely, the absence of key metabolic genes as phylogenetic signal underscores the utility of free-living flatworms as model systems for the parasitic lineages and the development of anti-helmith reagents.

A MADI–MAD2–independent spindle check–point? The apparent absence of MADI and MAD2 in planarians (Fig. 3c) raises the question of whether planarians have a functional SAC, and how essential cellular functions can be maintained in the absence of supposed core components. Both MADI and MAD2 (also known as MAD1LI and MAD2L1) are near-universally conserved owing to their essential roles in the SAC, which guards against aneuploidy by inhibiting cell cycle progression as long as even a single chromosome remains unattached to the mitotic spindle.
Although MAD1 and MAD2 homologues are easily identifiable in all other flatworms examined (Extended Data Figs 7, 8), not even flatworm queries could identify significant homologues in S. mediterranea or the transcriptomes of five other planarian species. Therefore, planarians are likely to have lost MAD1, MAD2 and multiple other SAC components (Fig. 4a). The known M-phase arrest of planarian cells upon pharmacological interference with spindle function (Fig. 4b) is therefore remarkable, as it indicates the maintenance of a SAC-like response despite a lack of supposed SAC core components.

To explore the underlying mechanisms of the SAC-like response in S. mediterranea, we targeted remaining components of the SAC network (Fig. 4a) by RNA interference (RNAi) and quantified the fraction of M-phase arrested cells with or without the microtubule depolymerizing drug nocodazole (Fig. 4b, Supplementary Information S18). The marked increase in the proportion of M-phase cells and the subsequent loss of dividing cells under RNAi targeting CDC20 (Fig. 4b, Extended Data Fig. 9a) or the anaphase-promoting complex/cyclosome (APC/C) subunit gene CDC23 (Fig. 4b) is therefore remarkable, as it indicates the maintenance of a SAC-like response despite a lack of supposed SAC core components.

The resulting bird’s eye view of a ‘difficult’ genome using long-read sequencing and de novo assembly also highlights important challenges that remain to be overcome. In the case of S. mediterranea, these include an abundance of low-complexity microsatellite repeats, inbreeding resistant heterozygosity and a new class of extraordinarily long LTR elements. However, the fact that the scaffold size of newly reported genome assemblies often remains substantially below the 3.85 Mb of the S. mediterranea assembly (Extended Data Table 1) indicates that similar genomics improvements of the MARVEL assembler towards heterozygous and/or compositionally biased sequencing data15 will be useful for enhancing assembly contiguity in de novo genome sequencing projects.

Discussion
We have described the highly contiguous genome sequence of the planarian model species S. mediterranea, which enables the genomic analysis of whole-body regeneration, stem cell pluripotency, lack of organismal ageing and other notable features of this model system.
We have also found a high degree of structural rearrangement and the absence of a number of conserved genes in the *S. mediterranea* genome. However, *D. melanogaster*, *C. elegans* and other animals also show loss of 'essential' genes\(^{15,26,32}\), which raises a general conundrum: how can animals survive and compete while lacking core components of essential mechanisms? In cell biological terminology, a core mechanism signifies a chain of molecular interactions that explain a given process in multiple species, while essentiality indicates importance for organismal survival. The emergence of viable yeast strains upon deletion of essential genes\(^{35}\) or the competitiveness of hundreds of extant planarian species in a diversity of habitats worldwide\(^{34}\) both make it clear that essentiality is relative. The demonstration of SAC function in the likely absence of MAD1 and MAD2 suggests that our genetic and mechanistic understanding of SAC function is incomplete. Further studies on planarians and other ‘non-traditional’ model organisms are needed to understand the basis and mechanism of these cellular functions. Such a function-oriented, rather than gene-centric, view of biological mechanisms abstracts general function from individual molecules and is therefore likely to ultimately facilitate the reverse engineering of biology.

**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

**Data Availability** The *S. mediterranea* genome assembly is accessible at GenBank under accession number NRG000000000 and can also be browsed at and downloaded from http://planmine.mpi-cbg.de. All DNA and RNA reads were deposited at the Sequence Read Archive under the bioproject accession PRJNA379262 and under the following SRA accession numbers: PacBio P4/C2 data, SRX2700681 and SRX2700682; PacBio P6/C4 data, SRX2700683; PacBio CCS data, SRX2700684; DNA shotgun, SRX2700686; DNA Chicago, SRX2700687; and RNA-seq, SRX2700685.

**Code Availability** The current source code of the MARVEL assembler is available at https://github.com/niolab/MARVEL. The execution scripts used for *S. mediterranea* can be found in the smed subfolder of the examples folder.

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**Figure 4 | Spindle assembly checkpoint (SAC) function in the likely absence of MAD1–MAD2.** a, Cartoon illustration of SAC core components and function. Black and red denote components conserved or missing in *S. mediterranea*, respectively. KMN network: KNL1, MIS12 complex, NDC80 complex. b, Fractional abundance of mitotic cells under RNAi targeting the indicated SAC component genes, with (red) and without (cyan) nocodazole pre-treatment. Values are shown as mean with 95% confidence intervals (n = 4 biological replicates, 10 pooled animals, 5 technical replicates with 5 or 6 images each). Cells treated with RNAi targeting CDC20 are shown as single replicates owing to rapid stem cell loss (Supplementary Information S18, Extended Data Fig. 9a, b). Significance assessed by two-way ANOVA, followed by Dunnett’s post-hoc test (****P < 0.0001; NS, not significant), excluding RNAi targeting CDC20.
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**Supplementary Information** is available in the online version of the paper.

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Extended Data Figure 1 | See next page for caption.
Extended Data Figure 1 | *S. mediterranea* sequencing and assembly quality control. a, *S. mediterranea* genomic DNA preparation. The established protocol (top) yields a black solution owing to co-purification of porphyrin pigments. The improved protocol (bottom) removes contaminants including the pigment and therefore results in clear preparations. b, The improved protocol consistently yields high-molecular-weight DNA, as shown by the pulse field gel electrophoresis of two independent preparations (right-hand two lanes) and DNA size markers (left-hand two lanes). c, Overview of all PacBio sequencing runs for the *S. mediterranea* assembly. d, Sequencing statistics of a representative PacBio RS II SMRT cell (P6/C4 chemistry). Total output: 1,053.4 Mb; reads of insert: 976.4 Mb; maximal read length: 52,441 bp. e, Connectivity matrix plot illustrating Chicago library read-pair distances after HiRise scaffolding. Colour coding identifies individual contigs contributing to the scaffold dd_Smed_{g4}_1. f, Mapping characteristics of *S. mediterranea* transcriptomes against the genome assembly with more than 60% query coverage and more than 60% sequence identity as cut-off criteria. Left, the dd_Smes_v1.PCFL transcriptome of the sequenced strain. Right, the dd_Smed_v6.PCFL transcriptome of the asexual strain. The pie charts show the absolute number and relative proportions of transcripts mapping with the indicated characteristics. g, Further analysis of the 538 non-mapping *S. mediterranea* transcripts from e (Supplementary Information S7). Missing gene, transcripts that map uniquely to the SmedSxl v4.0 assembly10 and have annotated orthologues in at least five other planarian species in PlanMine18. Putative contaminant, Top RefSeq BLAST hit in a likely contaminant species. Unknown, all remaining transcripts. The fact that only 46 out of 31,966 *S. mediterranea* transcripts are classified as genuinely missing indicates that the *S. mediterranea* assembly is largely complete. In contrast, 1,229 transcripts that uniquely mapped to the *S. mediterranea* genome and had orthologues in at least five other planarian species failed to map to the previously published SmedSxl v4.0 assembly10. Substantial gaps in the previous assembly also mean that the number of missing genes in the *S. mediterranea* assembly may be slightly higher, as some may have been classified as unknown.
Extended Data Figure 2 | See next page for caption.
Extended Data Figure 2 | Assembly validation by high stringency transcript back-mapping. a, Quality control of the *S. mediterranea* assembly by means of high stringency back-mapping of 1,509 high confidence (HC)-cDNAs. HC-cDNAs were defined as having BLAST hits with more than 90% query and subject coverage in seven other planarian transcriptomes in PlanMine.16 HC-cDNAs were mapped to the *S. mediterranea* assembly using more than 90% query coverage and sequence identity as cut-off criteria. The pie chart shows the absolute number and relative proportions of HC-cDNAs mapping with the indicated characteristics. b, Further analysis of the ten HC-cDNAs classified as non-mapping from a by intersection with the mapping results of Extended Data Fig. 1g. Of these, two were designated as ‘false negative’, as both mapped to the *S. mediterranea* genome with more than 90% query coverage and sequence identity using BLAT. c, UCSC genome browser screenshot (75-kb window) of the genomic mapping location of one of the two ‘unknown’ HC-cDNAs as a single example of a mapping failure due to an actual assembly error. The example documents inversion of the 5′-end of the cDNA within a low-confidence stretch at a contig end (lack of coverage in the Quiver track). The inversion is supported by inverted RNA-seq read mapping and inversion of the cDNA sequence shown in the respective tracks. Below, colour-coded Miropeats similarity plots of respective regions. d, e, Examples of genomic mapping loci of HC-cDNA transcripts from the multi-mapping category in a, browser screen shots as described in c. d, Example of a likely legitimate (biological) gene duplication in a gap-free high-confidence region. e, Micro tandem duplication surrounding a scaffolding gap in a repeat-rich region. f, Multi-mapping HC-cDNAs map preferentially to contig ends. The histogram plots the distance to the closest gap or contig end for the 67 multi-mappers and a corresponding number of unique mappers (a). g, Estimated size of the duplicated regions of multi-mapping HC-cDNAs. This analysis identifies a small fraction of small-scale duplications at assembly gaps in the *S. mediterranea* assembly, which can be easily identified with the help of the various quality control tracks in the PlanMine genome browser.
Extended Data Figure 3 | Repeats in the  *S. mediterranea* assembly.

*Panel a.* Estimation of the abundance of solo and full-length LTR elements in the *S. mediterranea* assembly. Elements SLF-8 and SLF-14 show a large number of solo LTRs compared to full-length copies, indicating a large number of excision events by homologous recombination. Of the Burro elements, Burro-1 was the most abundant, with 124 full-length copies, followed by Burro-3 and Burro-2 with 25 and 23 full-length copies, respectively.

*Panel b.* Comparison of lengths of indicated repeat consensus classes in *H. sapiens*, *D. melanogaster* and *C. elegans*. For *S. mediterranea*, we used a custom library generated in this study. Dark colours indicate predominant lengths of specific repeat classes. Red, repeat consensuses more than 15 kb in length.

*Panel c.* Expression analysis of gypsy LTR elements in *S. mediterranea* RNA-seq data using TETranscripts. The three most transcriptionally active elements were Burro-1, Burro-2 and SLF-8.

*Panel d.* LTR expression analysis by whole-mount in situ hybridization and single-cell expression data. Top, SLF-9-derived transcript. Bottom, Burro-1-derived transcript. Both are broadly transcribed in many *S. mediterranea* cell types (CIW4 strain, *n* = 1 biological replicate, 10 animals). Scale bar, 250 µm.

*Panel e.* Kimura distance plot of *S. mediterranea* LTR elements. Substitution levels varied by element, but also within element groups. Burro-1, Burro-2, Burro-3 and SLF-8 all contain elements spread over a large range of substitution levels, possibly indicative of continued activity over large time scales. The remaining elements are characterized by more defined peaks in expansion, with the highest average divergences being seen in the smallest elements characterized (SLF-10, SLF-11, SLF-12), making these amongst the oldest within the genome. Notably, both SLF-8 and SLF-9 have representative elements with particularly low substitution rates, potentially indicating a recent or ongoing expansion.
Extended Data Figure 4 | AT-rich microsatellites in the *S. mediterranea* genome. **a**, Features of AT-rich microsatellites. Top, inter-repeat spacing of repeats more than 99 bp in length. Bottom, repeat length. AT-rich microsatellites with an average length of 127 bp occur every approximately 3,500 bp. **b**, Genomic distribution of repeats more than 99 bp in length. **c**, Increased probability of read alignment termination within microsatellite repeats. Individual size bins were analysed separately for microsatellite repeats (red) or non-repetitive regions (cyan). Although accounting for only 4.2% of the assembly size, microsatellite repeats significantly limit assembly contiguity owing to an increased probability of read alignment loss. **d**, Genome-wide coverage ratios of insertion or deletion sequences more than 99 bp in length and excluding AT repeats. **e**, Read length variation analysis across AT-rich repeat regions (AAT) in regular PacBio sequencing data compared to circular consensus sequencing (CCS) coverage of the same region. CCS reads sample the same genomic region multiple times. The lack of a clear difference in the length variation of specific AT repeats between repetitive sequencing of the same DNA molecule (CCS) and that of sequencing reads representing different DNA molecules (regular PacBio data, P6/C4) indicates that repeat length variations are mainly technical in nature. Rather than repeat length polymorphisms, the most likely cause of the detrimental effect of the repeats is the increased ambiguity in low complexity sequence alignments (Supplementary Information S11.4). Unique (UQ) regions were included as controls. CCS_UQ: CCS subread length variation versus the consensus length of all subreads in binned unique regions (n = 3,300). CCS_AAT: CCS subread length variation versus the consensus length of all subreads in binned AT repeat regions (n = 4,825). P6_UQ: Length variation of individual reads in the regular PacBio sequencing data (P6/C4) versus the consensus length of the region in the *S. mediterranea* assembly in binned unique regions (n = 3,310). P6_AAT: Length variation of individual reads in the regular PacBio sequencing data (P6/C4) versus the consensus length of the region in the *S. mediterranea* assembly in binned AT repeat regions (n = 5,085). Dots, outliers; central horizontal line, median; box ranges, from first quartile to third quartile; whiskers, interquartile range (midspread): 75th and 25th percentile.
Extended Data Figure 5 | See next page for caption.
Extended Data Figure 5 | Comparative genomics. a, Contig and scaffold N50 statistics of the genomes used for the comparative genome alignments in Fig. 3b. The basal vertebrate lamprey genome assembly is more fragmented (similar or lower N50 values) than most other platyhelminth genomes. Nevertheless, the human-to-lamprey genome alignment has equivalent or even higher alignment chain scores and span lengths than any of the platyhelminth genome alignment comparisons, indicating that the true extent of sequence divergence and loss of conserved gene order in platyhelminths is likely to be greater than estimated. b, Example of a top-scoring alignment chain. The UCSC genome browser screenshot of the *S. mediterranea* genome shows that alignments predominantly overlap exons of the two transcripts shown at the top. This example is one of the few cases of apparent gene order conservation between *S. mediterranea* and *S. mansoni*. Blocks in the alignment chains represent local alignments, connecting single lines represent deletions in the query genome and double lines represent regions with sequence in both *S. mediterranea* and the query genome that do not align. c, Comparative loss analysis of highly conserved genes across the 26 indicated species. Red, conserved gene fraction, defined as the proportion of orthogroups containing at least 9 out of the 14 non-flatworm species and the query species. Blue, lost fraction of highly conserved genes, defined as the proportion of orthogroups containing at least 9 out of the 14 non-flatworm species, but not the query species (Supplementary Information S17). Absolute numbers of highly conserved genes are shown at the top, with slight fluctuations caused by species-specific sequence duplications.
Extended Data Figure 6 | See next page for caption.
Extended Data Figure 6 | Planarian-specific genes. a, Conservation of 1,165 flatworm-specific genes (Supplementary Information S16.1) amongst flatworm species. Only 61 sequences had sequence homologues in the indicated flatworm species (other denotes *Taenia solium, Echinococcus multilocularis, Echinococcus granulosus, Hymenolepis microstoma*), indicating that this gene set mostly represents planarian-specific genes. b, c, Characteristics of planarian-specific genes. b, Distribution of exon numbers compared to a control gene set (HC-cDNAs; Extended Data Fig. 2a), indicating enrichment of single-exon genes. c, Number of predicted domains (InterProScan), indicating that only a minority of genes contain predicted domains. d, Identity of detected domains (Pfam and SUPERFAMILY). Unintegrated signatures, recurring sequence motifs that are not grouped into InterPro entries. These might represent so far uncurated or weakly supported motifs that do not pass InterPro’s integration standards. e, Differential expression of 626 planarian-specific genes in published *S. mediterranea* RNA-seq data sets of different regeneration phases (left), stem cells or progeny populations (middle) or specific developmental stages (right). Red lines indicate differential expression relative to the control of each series (white indicates no change). Genes were ordered using rank by sum. The high proportion of differential expression indicates the widespread contribution of lineage-specific genes to planarian biology. f, g, Specific examples of non-conserved genes. Top, SMART domain representation. Bottom, Differential expression under the indicated conditions.
Extended Data Figure 7 | Sequence conservation of MAD1 protein in non-planarian flatworms. a. COBALT multiple protein sequence alignment of the MAD1 homologues of the indicated species (including all the non-planarian flatworm species from Fig. 3c). b. Heat map of BLOSUM62 sequence similarity matrix generated from alignment in a, demonstrating significant sequence conservation of MAD1 homologues even in flatworms.
Extended Data Figure 8 | Sequence conservation of MAD2 protein in non-planarian flatworms. a, COBALT multiple protein sequence alignment of the MAD2 homologues of the indicated species (including all the non-planarian flatworm species from Fig. 3c). b, Heat map of BLOSUM62 sequence similarity matrix generated from alignment in a), demonstrating significant sequence conservation of MAD2 homologues even in flatworms.
Extended Data Figure 9 | Effect of RNAi targeting \textit{CDC20} and SAC components on the planarian stem cell compartment. a, Fluorescent whole-mount \textit{in situ} hybridization of the planarian head region. Stem cells (neoblasts) were visualized using a \textit{smedwi-1} probe (red), early and late progeny by pooled \textit{prog-1} and \textit{agat-1} probes (green). Blue, nuclear counterstaining by DAPI. Top, RNAi control against \textit{EGFP}; bottom, RNAi targeting \textit{CDC20}, which results in a markedly decreased number of \textit{smedwi-1} and \textit{prog-1/agat-1} positive cells after three rounds of RNAi feeding. This indicates the loss of neoblasts and a concomitant reduction in progenitor numbers ($n = 1$ biological replicate, 10 animals). Scale bar, 200\,$\mu$m. b, Effect of indicated RNAi treatments on planarian stem cell abundance. Representative images of cell macerates, stained with DAPI (nuclei, blue), anti-H3ser10P (mitotic cells, magenta) and \textit{smedwi-1} \textit{in situ} hybridization (stem cells, yellow). Numbers indicate the mean fraction \( \pm \) s.d. of \textit{smedwi-1} positive cells of total cells quantified by nuclear counting using DAPI ($n = 1$, 10 pooled animals, 5 technical replicates with 5 images each). Scale bar, 50\,$\mu$m.
Extended Data Table 1 | *S. mediterranea* genome assembly comparisons

| Assembly       | SmedSkl v4.0   | GCA_000691995.1 | PacBio-Casu | PacBio-MARVEL | g4 assembly       |
|----------------|----------------|-----------------|-------------|---------------|-------------------|
| Technology     | Sanger         | Illumina        | PacBio      | PacBio        | PacBio + Chicago  |
| Assembler      | NA             | SOAPdenovo      | Camo        | MARVEL        | MARVEL + HiRise   |
| Assembly length (Mb) | 787.5          | 700.7           | 938.8       | 782.1         | 774.0             |
| **Contigs**    |                |                 |             |               |                   |
| # contigs      | 112,641        | 108,794         | 7,637       | 1,839         | 1,292             |
| Longest contig | 149,108        | 132,670         | 2,212,985   | 4,363,926     | 5,343,607         |
| Contig N50     | 11,977         | 10,721          | 194,023     | 788,691       | 1,121,568         |
| **Scaffolds**  |                |                 |             |               |                   |
| # scaffolds    | 15,134         | 12,782          | NA          | NA            | 481               |
| Longest scaffold | 893,023       | 1,050,243       | NA          | NA            | 17,761,579        |
| Scaffold N50   | 80,447         | 83,932          | NA          | NA            | 3,854,848         |
| % in gaps      | 13.97          | 14.32           | 0           | 0             | 0.01              |

Final Smed_g4 assembly characteristics are shown in bold.
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Nature Research wishes to improve the reproducibility of the work that we publish. This form is intended for publication with all accepted life science papers and provides structure for consistency and transparency in reporting. Every life science submission will use this form; some list items might not apply to an individual manuscript, but all fields must be completed for clarity.

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Experimental design

1. Sample size
   Describe how sample size was determined.
   All reported experiments used at least 6 animals. No statistical methods were used to determine sample sizes.

2. Data exclusions
   Describe any data exclusions.
   No data was excluded.

3. Replication
   Describe whether the experimental findings were reliably reproduced.
   All experimental manipulations were carried out at least in triplicate. The only exception is the cdc20RNAi point in Figure 4B/C. The main point of this figure is that stem cells become rapidly depleted upon cdc20(RNAi). The very few remaining stem cells in the 1 replicate shown are supported by no remaining stem cells in replicates 2 and 3 (not shown), therefore also this conclusion is supported by multiple replicates.

4. Randomization
   Describe how samples/organisms/participants were allocated into experimental groups.
   Experimental animals were randomly selected from large laboratory populations.

5. Blinding
   Describe whether the investigators were blinded to group allocation during data collection and/or analysis.
   Blinding was not used in this study. However, the results in Fig. 4 are based on validated automated quantification routines, which rule out observer bias.

Note: all studies involving animals and/or human research participants must disclose whether blinding and randomization were used.

6. Statistical parameters
   For all figures and tables that use statistical methods, confirm that the following items are present in relevant figure legends (or in the Methods section if additional space is needed).
   Confirmed
   - The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement (animals, litters, cultures, etc.)
   - A description of how samples were collected, noting whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
   - A statement indicating how many times each experiment was replicated
   - The statistical test(s) used and whether they are one- or two-sided (note: only common tests should be described solely by name; more complex techniques should be described in the Methods section)
   - A description of any assumptions or corrections, such as an adjustment for multiple comparisons
   - The test results (e.g. P values) given as exact values whenever possible and with confidence intervals noted
   - A clear description of statistics including central tendency (e.g. median, mean) and variation (e.g. standard deviation, interquartile range)
   - Clearly defined error bars

See the web collection on statistics for biologists for further resources and guidance.
Software

Policy information about availability of computer code

7. Software

Describe the software used to analyze the data in this study.

The assembler (Marvel) has been deposited in a GitHub repository. All other software packages and tools cited in the manuscript are referenced (either in main text or Supplemental Online Material section) and, where applicable, parameter settings are listed in the Supplemental Online Material section.

For manuscripts utilizing custom algorithms or software that are central to the paper but not yet described in the published literature, software must be made available to editors and reviewers upon request. We strongly encourage code deposition in a community repository (e.g. GitHub). Nature Methods guidance for providing algorithms and software for publication provides further information on this topic.

Materials and reagents

Policy information about availability of materials

8. Materials availability

Indicate whether there are restrictions on availability of unique materials or if these materials are only available for distribution by a for-profit company.

All material are available upon request from the Rink lab.

9. Antibodies

Describe the antibodies used and how they were validated for use in the system under study (i.e. assay and species).

Primary antibodies: anti-Histone H3 antibody (rabbit, phospho S10 + T11) [E173]; Abcam, Cat. No.: ab32107
Secondary antibodies: goat anti-rabbit IgG (H+L), Alexa Fluor 647; ThermoFisher Scientific, Cat. No.: A-21245
The primary antibody is an established tool in the field, as corroborated by the reference cited in the Supplemental Online Material section.

10. Eukaryotic cell lines

a. State the source of each eukaryotic cell line used.

b. Describe the method of cell line authentication used.

c. Report whether the cell lines were tested for mycoplasma contamination.

d. If any of the cell lines used are listed in the database of commonly misidentified cell lines maintained by ICLAC, provide a scientific rationale for their use.

Does not apply to this study.

Does not apply to this study.

Does not apply to this study.

Does not apply to this study.

Animals and human research participants

Policy information about studies involving animals; when reporting animal research, follow the ARRIVE guidelines

11. Description of research animals

Provide details on animals and/or animal-derived materials used in the study.

Individuals of clonal laboratory strains of the sexual and asexual lines of S. mediterranea were used in this study. Size and or feeding history of experimental specimens are listed as physiologically meaningful designators of experimental specimens.

Research on planarians is not subject to any ethical regulations in Germany.

Policy information about studies involving human research participants

12. Description of human research participants

Describe the covariate-relevant population characteristics of the human research participants.

Does not apply to this study.