Brief Communication

CRISPR-Cereal: a guide RNA design tool integrating regulome and genomic variation for wheat, maize and rice

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The clustered regularly interspaced short palindromic repeat (CRISPR)-associated protein (Cas) genome editing system (CRISPR-Cas) is revolutionizing agriculture. In this system, a guide sequence that matches to a particular genomic DNA is placed in front of a synthetic RNA that consists of a scaffold sequence necessary for Cas-binding to form a guide RNA (gRNA). gRNA/Cas complex binds to the target DNA that contains a protospacer adjacent motif (PAM) via base-paring and generates a double-strand break (DSB) by Cas protein. Mutations will be created when the DSB cannot be perfectly repaired. Among kinds of Cas nucleases with highest edit efficiency, NGG (N = A, T, G or C) for O. sativa cv Nipponbare (IRGSP-1.0) and the B73 maize genome AGPv4 (Jiao et al., 2017) are used as reference genomes. Notably, the MHRS3 genome is the first gap-free genome in crops, which allows gRNA design for ‘dark area’ like centromeric region (Figure 1b). The gap-free genome MHRS3 unmasks more than 395 non-TE genes located in centromere regions (Song et al., 2021).

Previously, we generated a whole-genomic pool for scanning gRNA in maize (http://crispr.hzau.edu.cn/CRISPR-Local/), but this approach is not suitable for wheat which has a 16 Gb genome (Figure 1b), seven times bigger than maize (Appels et al., 2018; Jiao et al., 2017). It is an obstacle to speedy screen genome-wide off-targets in wheat. To solve the problem, CRISPR-Cereal applies the command-line tool, FlashFry to perform genome-wide scan for off-targets (McKenna and Shendure, 2018). FlashFry uses guide-to-genome aggregation model to scan the genome and supports screening for unconstrained number of mismatches for putative off-targets. Given that DNA cleavage by Cas9 allows three to five mismatches, we made the option of mismatches range from zero to five. CRISPR-Cereal could identify all the off-targets with less than four mismatches for each candidate guide in 49 s in wheat, 35 s in maize and 9 s in rice, which outperforms all the other gRNA design tools including E-CRISP (http://www.e-crisp.org/E-CRISP/), CRISP direct (http://crispr.dbcls.jp/) and wheatCRISPR (https://crispr.bioinfo.nrc.ca/WheatCrispr/) in which either only stands or cannot search for genome-wide off-targets speedily. The output of CRISPR-Cereal contains information for all off-targets, which can be downloaded for further comparison. We further compared the off-targets between CRISPR-Cereal and Cas-OFFinder (http://www.rgenome.net/cas-offinder/), and observed that the off-targets detected by the above two tools are very similar (details in help page of the website). Significantly, this is the first time to perform genome-wide off-target scan in wheat. We randomly submitted three different wheat DNA sequences in FASTA format to search genome-wide off-targets with 0–5 mismatches. It shows that the off-target numbers could reach up to several millions (Figure 1c), implying the importance of whole-genome scan for off-targets. The on-target and off-target scores are predicted with the widely used on-target metrics and cutting-frequency determination (CFD) scoring scheme (Doench et al., 2014, 2016). CRISPR-Cereal provides the information of GC content, position, proximal gene, location of gene structure element (promoter, exon, intron or intergenic), efficiency score of on-targets and genome-wide off-targets...
The workflow of gRNA design using CRISPR-Cereal.

The correlation between mutant frequency and chromatin accessibility in gRNAs target sites. (j) The SNP information on TG7 target region. (k) The genomes included in CRISPR-Cereal. (e) The regulome visualization page using GBrowse. (f) The regulome information in 2 kb region around the genomes. (c) The numbers of genome-wide off-targets are increased when allowed mismatches are from zero to five. (d) CRISPR-Cereal tool.

For the off-targets in wheat, CRISPR-Cereal specifies whether they belong to the homologous group in the A, B or D sub-genomes (Figure 1d), and telling users whether the selected gRNAs would cause unintended editing in the homologous genes from the sub-genomes.
to assist gRNA design (Figure 1d–f). We collected data sets of assay for transposase-accessible chromatin using sequencing (ATAC-seq), DNasel-hypersensitivity sequencing (DNasel-seq), and formaldehyde-assisted isolation of regulatory elements by sequencing (FAIRE-Seq) to locate open chromatin, data from chromatin immunoprecipitation sequencing (ChIP-seq) to present DNA methylation level. Information of the data sets was listed in http://crispr.hzau.edu.cn/CRISPR-Cereal/help.php. The data sets were reanalysed and could be easily visualized by Generic Genome Browser (GBrowse) 2.0 (https://github.com/GMOD/GBrowse) after clicking the on-target position (e.g. 1A:-1264382) on the elementary result page of CRISPR-Cereal (Figure 1d,e). Users could extend to a widely region to see the global regulome information around targets (Figure 1f). Besides, to further help users decide which gRNA to use, the chromatin status of the targets has been scored (Figure 1d). The chromatin accessibility information helps to choose and design gRNAs. Recently, Gong and colleagues reported that when gRNA targeted 350 bp upstream of OsWOX11, the edit efficiency was high (Gong et al., 2020). We found that the reason might be due to the open chromatin feature at that region (Figure 1g). In addition, the gRNA for TG2 that failed to activate transcription is partly located in an un-open chromatin region (Gong et al., 2020) (Figure 1h). To further confirm the relationship between editing efficiency and chromatin accessibility on the gRNAs target sites, we randomly collected the published data for 84 endogenous sites and checked the chromatin accessibility on the corresponding on-target sites in rice callus (Zhang et al., 2012). As expected, gRNAs targeting open chromatin regions result in significantly higher editing efficiency than those against un-open regions (R = 0.34, P = 0.002) (Figure 1i).

To expand the application of CRISPR-Cas tool from reference genome to elite cultivars, CRISPR-Cereal collected and visualized SNPs information from Ensembl Plants (ftp://ftp.ensemblgenomes.org/pub/plants/release-48/variation/vcf/) and RiceVarMap v2.0 (http://ricevarmap.ncpr.cn/) for rice, MaizeSNPDB (https://venya.oxzy/MaizeSNPDB/) for maize and 487 wheat genotypes (Pont et al., 2019). Remarkably, we found that 39.59% guide sequences in rice genome possess SNPs, which would cause mismatches if guide sequence would be decided only by reference genome. One example is that the gRNA for TG7 designed based on the Nipponbare genome has 6 SNPs failed to mediate transcription activation of OsWOX11 (Gong et al., 2020) (Figure 1j), although the location and chromatin openness may also play roles in this process.

In summary, CRISPR-Cereal integrates regulome information and considers SNPs existed in the candidate gRNAs to promote precise and high-efficient gene editing for wheat, maize and rice. The workflow of CRISPR-Cereal is shown in Figure 1k, it is freely available at http://crispr.hzau.edu.cn/CRISPR-Cereal/.

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Conflict of Interest

The authors declare no conflict of interest.

Author contributions

C.H. and H.L. collected the data and built the gRNA design platform and GBrowse visualization platform. C.H. drafted the manuscript with input from L.-L.C. and W.Y. D.C. analysed the histone modifications data of wheat and rice. W.-Z.X. analysed the genomic variation data of rice. M.W., Y.L. and X.G. contribute to generate the web-page. L.-L.C. and W.Y. conceived of the study and wrote the manuscript. All authors read and approved the final manuscript.

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