Full Paper

Improvement of barley genome annotations by deciphering the Haruna Nijo genome

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Edited by Dr Masahiro Yano
Received 3 August 2015; Accepted 26 October 2015

Abstract
Full-length (FL) cDNA sequences provide the most reliable evidence for the presence of genes in genomes. In this report, detailed gene structures of barley, whole genome shotgun (WGS) and additional transcript data of the cultivar Haruna Nijo were quality controlled and compared with the published Morex genome information. Haruna Nijo scaffolds have longer total sequence length with much higher N50 and fewer sequences than those in Morex WGS contigs. The longer Haruna Nijo scaffolds provided efficient FLcDNA mapping, resulting in high coverage and detection of the transcription start sites. In combination with FLcDNAs and RNA-Seq data from four different tissue samples of Haruna Nijo, we identified 51,249 gene models on 30,606 loci. Overall sequence similarity between Haruna Nijo and Morex genome was 95.99%, while that of exon regions was higher (99.71%). These sequence and annotation data of Haruna Nijo are combined with Morex genome data and released from a genome browser. The genome sequence of Haruna Nijo may provide detailed gene structures in addition to the current Morex barley genome information.

Key words: genome sequencing, full-length cDNA, Hordeum, RNA-Seq

1. Introduction
Barley (Hordeum vulgare L.) is used for many purposes including human food, malting and animal feed. Coupled with its wide adaptability in environments ranging from the highlands of Africa at the equator to the Arctic Regions in Scandinavia, barley is the fourth most important cereal crop (http://faostat.fao.org, 7 November 2015, date accessed). In addition, barley is an ancient crop that was domesticated ca. 10,000 yrs ago in the Fertile Crescent.1 Decades of collection and curation of barley germplasm have resulted in substantial germplasm collections including: IPK (Germany), USDA-ARS (USA) and Okayama University (Japan).2 To fully exploit these in situ and ex situ collections for barley breeding and gene discovery, barley genome sequences of multiple genotypes are needed.

To generate the first genome sequence of barley, the International Barley Sequencing Consortium (IBSC) conducted BAC fingerprinting, BAC end sequencing, whole genome shotgun (WGS) sequencing, RNA-Seq analysis and genetic mapping and integrated these data to develop a gene-based genome sequence of the North American six-row spring malting barley cultivar Morex.3 Morex traces to barley germplasm of Manchurian origin with six-row inflorescence, which is not common for malting barleys other than the USA. Currently, Morex is the reference genome for barley genetics studies, and additional efforts are in progress to improve the Morex genome sequence.4,5 Three other barley genomes including Bowman, Barke and a Tibetan hulless genotype have been WGS sequenced3,6 and provide additional sequence information for comparison to Morex.
Haruna Nijo is a Japanese malting barley cultivar exhibiting excellent malting quality for brewing beer. Historically, Japanese malting harleys have been developed by crossing European malting barley cultivars and Japanese landraces and selecting for malting quality and adaptation to Japanese environments. Haruna Nijo was also developed by this approach in 1979 and has been extensively used as a foundation genotype of current Japanese breeding. Haruna Nijo has a high thermostability of β-amylase, and it also shows different Bmy1 allele for β-amylase activity from Morex. It is also shown that Haruna Nijo has a high malt extract (related to beer production) by QTL analysis. To enhance the utility of Haruna Nijo for breeding, a suite of genomics resources were developed including: an EST collection, transcript map construction, a BAC library, 454-based WGS sequencing and full-length (FL) cDNA sequencing and analysis (available at http://barleyflc.dna.afrc.go.jp/bexdb/, 7 November 2015, date accessed, with functions of expression profiling and genome browsing).

In the case of barley genome resources, the Morex reference genome and the Haruna Nijo FLcDNA sequences provide unique resources in different genotypes. Thus, to enhance the utility of both resources they need to be compared and integrated. Sequence comparisons of the two different malting barley haplotypes will be much more useful than single haplotype information, as they provide the basis of structural and functional allelic diversity. Previous comparisons of WGS data and FLcDNA sequences from Haruna Nijo and the genome sequence from Morex demonstrated a large amount of sequence polymorphisms between the two genotypes. For example, the genetic distance (dissimilarity) between Morex and Haruna Nijo by 1,536 SNPs of Illumina GoldenGate Assay is 0.50, which is larger than the difference (0.48) between Morex and Akashinriki (Japanese food barley). Thus, the Haruna Nijo FLcDNA and WGS sequences may provide useful gene annotations for the reference genotype Morex and a resource for future breeding and gene discovery activities.

Our overall goal is to provide genome information of Haruna Nijo to annotate gene structures on the genome sequence of Morex. The four specific objectives of this study were to (i) generate high-quality genomic sequence of Haruna Nijo, (ii) map the precise position of genes identified in the Haruna Nijo genome, (iii) compare gene information between Haruna Nijo and Morex, and (iv) develop gene models based on the Haruna Nijo sequence data.

2. Materials and methods

2.1. Plant materials and nucleic acid isolation

For DNA and RNA isolation from seedling tissues, seeds of Haruna Nijo were germinated on moist filter paper in Petri dishes at 20°C in the dark. For DNA isolation, shoots of ca. 5 cm were harvested and the RNA samples from immature spike and immature seed samples, barley seeds were planted in pot with soil mixture (N: 120 mg l−1; P: 100 mg l−1; K: 160 mg l−1, pH 5.5) and grown in the greenhouse at 20/15°C day/night temperature under natural light conditions. Immature spike samples were harvested from the leaf sheath 5 days before heading. Immature seeds were sampled 35 days after flowering (soft dough stage). Total RNA from immature seeds was isolated by TRIzol® Reagent (Life Technologies, Japan) following the Plant RNA Isolation protocol. All RNA samples were purified by RNeasy Plant Mini Kit (QIAGEN K.K., Japan), and DNA was removed by RNase-Free DNase Set (QIAGEN K.K., Japan).

2.2. Library development

2.2.1. 454 long paired-end library

The high-quality genomic DNA was fragmented in 8 k and 20 k by the HydroShear (Digilab Inc., Holliston, MA, USA). The library was developed by a library preparation kit (Roche diagnostics, Japan) for long-paired-end libraries (8 k and 20 k). Circularization was performed according to the Roche Paired End Library Protocol, using Roche circularization adapters (Paired End Library Preparation Method Manual 20 kb and 8 kb Span; Roche Diagnostics, October 2009, Steps 3.1–3.7.3). Subsequently, the circularized fragments were fragmented again by nebulization to develop a sequencing library.

2.2.2. Illumina HiSeq sequencing library

DNA (2 μg) was fragmented by nebulization. Libraries were prepared according to the manufacturer’s instruction ‘Preparing Samples for Paired-End Sequencing, Part # 1005063 Rev. A June 2008’ of TruSeq DNA Sample Prep Kit (Illumina Japan). DNA fragments were size selected (500 bp) following the Low-Throughput Protocol of TruSeq DNA Sample Prep Kit (Illumina Japan). The quality of the library (fragment length distribution) was checked by Agilent Bioanalyzer High Sensitivity DNA Assay (Agilent Technologies, Japan) and KAPA Library Quantification Kit (KK4835, Kapa Biosystems, MA, USA).

2.2.3. MiSeq RNA sequencing library

Libraries for RNA-seq analysis were developed from each RNA sample using TruSeq RNA Sample Prep Kit V2 (Illumina Japan). The protocol of TruSeq RNA Sample Preparation V2 Guide (Illumina Japan) was used with modification of fragment isolation from agarose gel electrophoresis and elution in EB (elution buffer) following the method in the TruSeq Sample Preparation Guide (Illumina Japan). The library was quantified with Agilent High Sensitivity DNA Kit (Agilent Technologies Japan) and Qubit 2.0 Fluorometer (Life Technologies Japan).

2.3. Sequencing

2.3.1. 454 FLX Titanium platform

Each library was emulsion PCR amplified. The PCR-amplified fragments on beads were washed, and the bead number was counted using a Coulter Counter Z1 single threshold instrument (Beckman Coulter Japan). The appropriate number of beads was applied on a picotitre plate according to the manufacturer’s protocol. The FLX Titanium platform was used for sequencing (average read length 500 bp). The pyrosequencing reaction data were base-called to generate sff format files using the software installed on the analysis server of the 454 sequencer (Roche diagnostics, Japan).

2.3.2. Illumina HiSeq platform

The shotgun library was sequenced on an Illumina HiSeq 2000 to produce 2 x 101 paired-end reads. Raw data processing, base calling and quality control were performed with manufacture’s standard pipeline. The quality of the output sequences was inspected using the FastQC program (http://www.bioinformatics.bbsrc.ac.uk/projects/fastqc/, 7 November 2015, date accessed).

2.3.3. Illumina MiSeq RNA sequencing

The RNA-Seq library was sequenced with MiSeq Reagent Kit V3 (2 x 300 bp cycles) on MiSeq NGS system according to the MiSeq System
2.4. Assembly, sequence trimming and repeat masking

2.4.1. WGS assembly

Illumina PE reads, 454 single-end reads and 454 long paired-end reads were hybrid assembled by the de novo assembly algorithm of CLC Assembly Cell ver. 3.2.2 installed on a linux server with a main memory of 256 Gb.

2.4.2. Sequence trimming and repeat masking

Human, fungi (33 species) and microbial (2,777 species) nucleotide sequence data were obtained from NCBI. Each assembled contig sequence was queried on sequences of these species, and the contigs showing >80% identity and >50% of cumulative coverage by blast+16 were assumed as alien sequences against the barley genome. For repeat masking, fasta files of Triticaceae repeat sequence database (TREP Release 10: n = 1,717) were downloaded from GrainGenes (http://wheat.pw.usda.gov/TIMI/Repeat/). In addition, new repetitive elements were generated from the Haruna Nijo genome assembly by RepeatModeler (http://www.repeatmasker.org/RepeatModeler.html, 7 November 2015, date accessed). These two libraries were used for repeat masking the Haruna Nijo assembly by censor17 with -mode norm mode. For the analysis of repeat distribution, only results of repeat masking by the TREP library were used. For comparison, genomes of *Triticum aestivum*, *Triticum urartu* and *Aegilops tauschii* were repeat masked in a similar manner as the Haruna Nijo genome.

2.5. FLcDNA/RNA-Seq mapping

2.5.1. FLcDNA mapping

Of the two independent projects,13,14 5,006 FLcDNAs were retrieved from NCBI13 and 23,614 FLcDNAs14 were downloaded from the bex-db (http://barleyflc.dna.affrc.go.jp/bexdb/). Morex genome assembly (contigs) was retrieved from the PGSB PlantsDB database (http://pgsb.helmholtz-muenchen.de/plant/plantsdb.jsp, 7 November 2015, date accessed). FLcDNAs were mapped both on the Haruna Nijo scaffolds and on the Morex contigs using blast+ with parameters ‘-task blastn-evalue 0.01 -lcase_masking -num_descriptions 100’ and blast hits with >98% identity and >50% coverage were assumed as rRNAs. tRNA genes were predicted using the tRNAscan-SE ver. 1.3.1 program.30 Any tRNAs that were annotated as ‘possible pseudo-genes’ were not counted.

2.5.2. RNA-Seq analysis

After trimming of low-quality nucleotide and adapter sequences by Trimmomatic,23 reads resulting in rRNA sequences were discarded by Bowtie2.24 A set of programs, Bowtie2,24 TopHat225 and Cufflinks,26 were used to map reads on the Haruna Nijo scaffolds. Gene structures derived from FLcDNA mapping were utilized to develop consensus gene models of the four library reads. To predict ORFs on the gene structures, blastx searches were conducted to RefSeq and UniProtKB data sets. From the best hit of blastx results, the most reliable ORFs were determined with at least 70 amino acids in length. To assign gene function, InterProScan527 was conducted on the predicted ORFs. Based on the InterPro domain, GO terms were assigned to each ORF, and GOslim28 (http://agbase. msstate.edu/cgi-bin/tools/goslimviewer_select.pl, 7 November 2015, date accessed) was conducted.

Table 1. Sources of WGS sequencing data for Haruna Nijo

| Method            | Read                  | No. of reads | No. of bases (bp) |
|-------------------|-----------------------|--------------|-------------------|
| Illumina Paired end (PE) | 3,005,632,276            | 298,945,920,500 |
| 454 PE 20 kb      | 14,242,510              | 4,569,072,624  |
| 454 PE 8 kb       | 2,955,788               | 1,016,746,348  |
| 454 Single end 500 bp | 70,936,197             | 25,410,875,001 |

2.5.3. RNA gene prediction

From the silva database29 (http://www.arb-silva.de/, 7 November 2015, date accessed), 17 barley rRNA sequences were downloaded. They were mapped on the Haruna Nijo scaffolds using blast+ with the parameters ‘-task blastn-evalue 0.01 -lcase_masking -num_descriptions 100’ and blast hits with >98% identity and >50% coverage were assumed as rRNAs. tRNA genes were predicted using the tRNAseq-SE ver. 1.3.1 program. Any tRNAs that were annotated as ‘possible pseudo-genes’ were not counted.

2.5.4. Genome sequence comparison between Haruna Nijo and Morex

The Haruna Nijo scaffolds were mapped on the Morex contigs using megablast with the parameters ‘-evalue 1e-10 -num_descriptions 3’. Since several query sequences were hit on the same target sequence, the best hit regions were identified. If more than one Morex contig was positioned on a Haruna Nijo scaffold, these contigs were aligned on the scaffold allowing 10% overlap of the contig sequences.

2.5.5. Morex gene mapping

High confidence (HC) and low confidence (LC) genes of Morex were downloaded from the PGSB barley genome database and mapped to the Haruna Nijo scaffolds by GMAP31 with >95% identity and >90% coverage.

3. Results and discussion

3.1. Assembly of the Haruna Nijo genome sequence

We analysed different platforms of WGS sequences of Haruna Nijo by hybrid assembly. A total of 305 Gbp of sequence data were generated by Illumina HiSeq paired-end reads and 454 Titanium long paired-end reads (8 K and 20 K) (Table 1). The hybrid assembly of Illumina and 454 reads, which also include the published 454 Titanium single-end reads (25Gbp),3 generated a total of 2,055,601,874 bp in 1,753,384 scaffolds (contigs and gaps). After sequence trimming, the Haruna Nijo genome size was 2,005,970,762 bp in 1,712,236 scaffolds (contigs and gaps). The N50 was 3.5 kb, which is 2.5 times larger than the published Morex WGS contigs.3 The average sequence length in the published Morex WGS contigs was 1,3 kb. The Haruna Nijo scaffolds were longer than the Morex WGS contigs (Supplementary Fig. S1). Compared with Morex WGS contigs, which were assembled with only Illumina paired-end reads, the longer Haruna
Nijo WGS assembly is likely due to assembling the Illumina paired-end reads and 454 longer single-end reads.

3.2. Repeat content of the Haruna Nijo genome

Previous studies have shown that the barley genome contains ∼80% repeated elements. In the published WGS assemblies, the relative frequency was reduced to ∼60%, due to the degradation of redundant repeated sequences. On the assembly of Haruna Nijo WGS data, repeat analysis by the Triticeae repeat library (TREP) detected 55.9% as repeated sequences. After deleting unknown genomic regions (‘N’ sites), 60.8% of the assembled sequences were classified as repeated sequences. The composition and distribution of repeat sequence categories were similar between Haruna Nijo and Morex (Fig. 1). Noteworthy, Gypsy, Copia retrotransposons and the CACTA DNA transposon showed higher frequencies than those normally observed in other Triticeae species. When we compared repeat categories from Haruna Nijo, T. aestivum (common wheat) cv. Chinese Spring, T. urartu and A. tauschii, the frequency of Copia in Haruna Nijo was higher than those in the wheat genomes, while Haruna Nijo contained more CACTA and less Gypsy than those in wheat genomes (Supplementary Fig. S2).

To identify unique repeat sequences in barley, de novo repeat sequences in the Haruna Nijo scaffolds were detected by RepeatModeler, which identified a total of 979 consensus sequences. The de novo repeat library and the TREP library were used for repeat masking. As a result, 74.8% of the assembled sequences were masked (‘N’ sites were not masked and thus deleted from the calculation). The repeat masking by the de novo library identified 68.1% repeat sequences, which were much more than the masked repeats (60.8%) detected by the TREP library.

3.3. FLcDNA mapping

FLcDNA mapping on genome sequence provides reliable exon vs. intron structures on gene models. A non-redundant set of 28,620 FLcDNA sequences was mapped to the Haruna Nijo scaffolds. Blastn mapping revealed that 27,784 FLcDNAs (97%) showed sequence similarities with the Haruna Nijo scaffolds. According to the results of the FLcDNA mapping to scaffolds, we analysed mapping coverage of 5′-end of each FLcDNA on Haruna Nijo scaffolds. The ratio of completely mapped FLcDNAs, which were mapped from the first nucleotide sequence, was <10%. This shows that most of the FLcDNAs do not have the complete sequences on the 5′-side exon. To analyse additional mapped FLcDNAs, we relaxed the mapping position to the first 10 nucleotide sequences. Then, the ratio of mapped FLcDNA was raised to 80% (Supplementary Fig. S3). GC contents and CG-skew were analysed on the FLcDNAs having mapped 5′-ends within 10 bp. The typical high GC contents and CG-skewed peaks at the transcription start sites were identified as previously observed in rice and Arabidopsis (Supplementary Fig. S4).

The longer Haruna Nijo scaffolds may contribute to the efficiency in FLcDNA mapping. The number of mapped FLcDNAs under the threshold of 50% mapping coverage was 26,240 in the Haruna Nijo scaffolds and 24,261 in the Morex contigs (Table 3). These numbers included FLcDNAs that mapped to either the Haruna Nijo scaffolds (1,985) or the Morex contigs (379). However, if the threshold of mapping coverage was raised to 95%, the number of mapped FLcDNAs decreased to 14,044 on the Morex contigs, while 19,483 were still mapped on the Haruna Nijo scaffolds.

We also analysed the coverage of FLcDNA mapped reads on the Haruna Nijo scaffolds and the Morex contigs by comparing the mapped exon numbers of FLcDNAs between Haruna Nijo and Morex. Of the 24,255 FLcDNA gene models, 19,207 (79.2%) were mapped with the same number of exons both on the Haruna Nijo scaffolds and on the Morex contigs. Of these, 3,066 gene models had more exons in Haruna Nijo than that in Morex, while 1,982 gene models had more exons in Morex than Haruna Nijo. The start position analysis also showed that the Haruna Nijo scaffolds (6,842) had more mapped 5′-ends sequences than the Morex contigs (1,721).

These results indicated that the available genome sequences of Morex and Haruna Nijo are different, and there may be more cases...
where FLcDNAs mapped with higher coverage on Haruna Nijo genomes. For example, the entire FLcDNA (AK371953) for the *cleistogamy1* (*cly1* for closed flowering) gene, which was mapped on a Haruna Nijo scaffold, while the 5′ region was not mapped on a Morex contig (Fig. 2). Moreover, flanking regions for the FLcDNA (3.6 Kb for upstream and 2.9 Kb for downstream regions) were present on the Haruna Nijo scaffold to provide the opportunity to analyze the cis-element(s) of the gene.

### 3.4. Gene annotation

The IBSC presented 26,159 HC (high confidence) and 53,220 LC (low confidence) gene models by using RNA-Seq-based transcript and FLcDNAs. This number of genes was larger than that of the Haruna and 72.2% of Morex LC gene models on Haruna Nijo gene models.

| Coverage (%) | No. of hit FLcDNAs (ratio/total) |
|--------------|----------------------------------|
|              | Haruna Nijo | Morex[^3] |
| 50           | 26,241 (0.917) | 24,621 (0.860) |
| 70           | 22,835 (0.798) | 19,926 (0.696) |
| 80           | 21,429 (0.749) | 17,706 (0.619) |
| 90           | 20,153 (0.704) | 15,361 (0.537) |
| 95           | 19,483 (0.681) | 14,044 (0.491) |

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Figure 2. Gene structure of closed flowering locus *cly1* (AK371953). In the Haruna Nijo genome, the complete gene structures of *cly1* including upstream and downstream regions were identified. In the Morex genome,[^3] the first exon and upstream regions were truncated because of insufficient length of the contig.
the abundance of the retrotransposon Gypsy. The Haruna Nijo scaffolds also have higher rate of Gypsy; however, the number of tRNAs was smaller than that in wheat.

3.5. WGS assembly comparison between Haruna Nijo and Morex

There are several published and ongoing efforts of sequencing BACs derived from Morex. BAC-based assemblies may provide longer sequences than WGS assemblies to estimate gene models on the genome. The final goal for developing Haruna Nijo gene models is to map them on the BAC-based Morex genome sequences. To understand the quality of current Haruna Nijo genome resources, sequences of the Haruna Nijo scaffolds and the Morex contigs were aligned by megablast. An average sequence identity between Haruna Nijo and Morex was estimated as 95.99% (Supplementary Table S2). We found that mapped regions of query were different when query and database sequences were exchanged between Haruna Nijo and Morex. This was caused by the difference of sequence length in two assemblies. In this analysis, we used the best hit of megablast for the calculation, and the other hits with lower coverage on Haruna Nijo scaffolds were discarded. Therefore, the total length of aligned regions was decreased. Genome comparison of Haruna Nijo and Morex identified the occasions that the genomic regions of Haruna Nijo and Morex do not match. An example of mismatch between two assemblies on chromosome 5H was that a region of the Haruna Nijo gene model of TCONS_...
00033948 or TCONS_00033949 (Fig. 3A), which is supported by a
FLcDNA, AK370496 that encodes bZIP transcription factor family
protein on a contig_55003 (scaffold), was divided into two loci of
MLOC_14578.1 on morex_contig_156845 (Fig. 3B) and MLOC-
75543 on morex_contig_68185 (Fig. 3C). Thus, we may
find a better
gene model if we compare multiple gene models derived from different
assemblies.

We also analysed conserved genic regions between Haruna Nijo
and Morex by using mapped regions of FLcDNAs. The difference be-
tween Haruna Nijo scaffolds and FLcDNA sequences was 0.05%
(1 bp of 2000bp in exon), which is still five times higher than the
value of 0.01% in rice genome.37 However, the sequence difference
of exon regions between Haruna Nijo and Morex (identi-
fied by Har-
una Nijo FLcDNAs) was 0.29%, indicating that the sequence quality
of scaffolds in Haruna Nijo was high compared with the haplotype
difference between Haruna Nijo and Morex.

3.6. Conclusion
Based on this study, there are several advantages of using the Haruna
Nijo sequence resources. The Haruna Nijo scaffolds and FLcDNA sequences was 0.05%
(1 bp of 2000bp in exon), which is still five times higher than the
value of 0.01% in rice genome.37 However, the sequence difference
of exon regions between Haruna Nijo and Morex (identified by Har-
una Nijo FLcDNAs) was 0.29%, indicating that the sequence quality
of scaffolds in Haruna Nijo was high compared with the haplotype
difference between Haruna Nijo and Morex.

4. Availability
The short genomic reads used in the study are deposited at DDBJ-SRA
under accession ID PRJDB4103. The scaffolds for the Hordeum vul-
gare cv. Haruna Nijo are available at http://barleyflc.dna.affrc.go.jp/
bexdb/pages/harunanijo_index.jsp for download. Genome browsing
of the Haruna Nijo gene models are available on GBrowse [https://
gmod.org/wiki/GBrowse] at http://barleyflc.dna.affrc.go.jp/gb2/
gbrowse/HarunaNijo_genome/.21]. All scaffolds were concatenated
by 100 ‘Ns’ from the longer scaffolds. Haruna Nijo scaffolds,
FLcDNAs, RNA-Seq genes, repeat information, Morex contigs and
Morex HC/LC genes are displayed (Fig. 3). From the Haruna Nijo
genome browser, FLcDNA annotation data and Morex genome
browser of bex-db could be accessed (Fig. 3). Moreover, users can ac-
access the Haruna Nijo genome from FLcDNA information of bex-db.

Acknowledgements
We would like to express our thanks to Prof. Gary Muehlbauer, University of
Minnesota for his critical reading and linguistic editing of the manuscript. Bar-
ley seeds were provided through the National Bioresource Project of barley,
MENT of Japan. Analysis of Illumina HiSeq 2000 sequence reads was per-
formed by the Japan Advanced Plant Science Research Network (http://www.
psr-net.riken.jp/index.html). This work was partly supported by the scientific
technique research promotion program for agriculture, forestry fisheries and
food industry to K.S.

Supplementary data
Supplementary data are available at www.dnaresearch.oxfordjournals.org.

Funding
Funding to pay the Open Access publication charges for this article was pro-
vided by Okayama University to Kazuhiro Sato, corresponding author.

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