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Full Paper

Exploiting the great potential of Sequence Capture data by a new tool, SUPER-CAP

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Abstract

The recent development of Sequence Capture methodology represents a powerful strategy for enhancing data generation to assess genetic variation of targeted genomic regions. Here, we present SUPER-CAP, a bioinformatics web tool aimed at handling Sequence Capture data, fine calculating the allele frequency of variations and building genotype-specific sequence of captured genes. The dataset used to develop this in silico strategy consists of 378 loci and related regulative regions in a collection of 44 tomato landraces. About 14,000 high-quality variants were identified. The high depth (>40×) of coverage and adopting the correct filtering criteria allowed identification of about 4,000 rare variants and 10 genes with a different copy number variation. We also show that the tool is capable to reconstruct genotype-specific sequences for each genotype by using the detected variants. This allows evaluating the combined effect of multiple variants in the same protein. The architecture and functionality of SUPER-CAP makes the software appropriate for a broad set of analyses including SNP discovery and mining. Its functionality, together with the capability to process large data sets and efficient detection of sequence variation, makes SUPER-CAP a valuable bioinformatics tool for genomics and breeding purposes.

Key words: target enrichment, sequence reconstruction, heterozygous variants, web tool analysis

1. Introduction

Identifying sequence polymorphisms responsible for phenotypic variation and understanding the genetic basis of complex traits have been two major challenges of plant molecular genetics since the development of the first molecular markers up to the impressive highthroughput genomics technologies available today.^{1,2} The process of genotype-phenotype association is one of the central goals in the path towards plant improvement, and this requires the use of all the genetic and genomic information available for a given individual and/or population. The association of the genomic variation with traits of interest requires the reliable detection and the systematic investigation of the entire spectrum of DNA variability, including single nucleotide polymorphisms (SNPs), insertions/deletions (INDELs) as well as copy number variations (CNVs) and presence/absence of variations (PAV). Substantial progress towards this goal was made in the last few years. Large germoplasm collections have been characterized using SSR,^{3,4} SNP markers and SNP arrays.^{5–9} However, most of these approaches had known disadvantages that limited the

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power of the detection, in particular because the SNPs used in these studies were selected from a limited number of divergent sources and were commonly chosen to exceed a minimum frequency of the rare allele. Whole genome sequencing has also been used to explore individual variation at the genomic level in plants¹⁰⁻¹³ but, due to its high cost and complexity in data analysis, it is not expected to be widely applied for investigation of variants underlying specific traits of interest at a sufficient coverage. Recently, the development of microarray based or liquid-based genomic selection methods, commonly referred to as 'Sequence Capture', provided an affordable way to produce high-quality variants, thus solving most of the ascertained bias reported.¹⁴⁻¹⁶ Combining the recently developed targeted sequence enrichment with Next-Generation Sequencing (NGS) technologies, Sequence Capture methodology represents a powerful strategy for enhancing data generation to assess genetic variation of target regions, and it is likely to replace PCR as the main target enrichment method in both plants and animals.¹⁷ In addition, setting the experiments to obtain a sufficient depth of coverage would support the detection of rare functional variants and allow the discovery of complete haplotypes of genes, as well as CNV and PAV. With the ever-decreasing costs of sequencing and the advances in Sequence Capture technologies, these approaches are nowadays largely applied in different fields. Thanks to its high reliability and moderate cost per experiment, Sequence Capture is becoming an affordable diagnostic tool for medical and personalized medicine purposes.18 Despite its large employment in the medical field, the use of Sequence Capture in plant science is still emerging. Sequence Capture experiments were carried out for few species, including maize,19 strawberry,²⁰ rapeseed canola,^{21,22} black cottonwood,²³ potato,²⁴ wheat,25,26 medicago17 and cassava.27

The work herein proposed, which investigates the sequence variation of a group of 378 genes and the related regulative regions in a collection of 44 landraces, represents the first study of Sequence Capture in tomato species. Although software tools are available for variant calling and variant mining, their application is not straightforward, requiring users to install various packages and to convert data into different formats. This lack of easily accessible software pushed us to propose a web-based tool, named SUPER-CAP (http://supercap.sequentiabiotech. com/), to boost quick, proficient and affordable analysis of sequence capture data. This tool, benefitting from SUPER-W pipeline²⁸ and combining different software and customizable procedures, assists the user in sequence capture experiments from the identification of singlenucleotide variants (SNVs) and small insertions and deletions (INDELs) to the reconstruction of genotype-specific (hereafter referred to as 'private') sequences/gene features for each target region of each sample.

In addition, in this study, an explorative investigation on different aspects affecting sequence capture data was also carried out. For instance, since one of the major challenges in the enrichment/capture technologies are to avoid spurious variants calling for heterozygous (He) loci, this work evaluated how the depth of the reads and the procedure set-ups impact the variant calling of He variants. In fact, being the tomato a highly homozygous (Ho) species, it represents a good model for testing He variant calling into the gene set considered. In addition, spurious callings are frequently referred to multi-copy gene families (duplicated region/high homologous regions).²⁹ The gene set considered in this study represents a large range of variability in terms of gene families. This allowed for suitable assessment and estimation of the effective presence of He variants among the multi-copy genes.

As a final point, the variant set was also used to investigate: (i) the level and the type of sequence polymorphism in captured genes across the 44 tomato samples, (ii) the pattern of variants distribution among the lines, with respect to the identification of rare variants, (iii) the number of genes CNV, PAV and (iv) the functional annotation of the variants in order to prioritize the study of highly relevant variants. The work herein proposed represents a framework for easy Sequence Capture-related experiments that will promote similar studies, as well as in differing species.

2. Material and methods

2.1. Plant material and selection of target genes

A panel of 44 genetically diverse *Solanum lycopersicum* genotypes was selected from a wide collection of tomato landraces³⁰ available at the University of Naples, department of Agricultural Sciences. Plants were grown in a greenhouse under controlled conditions at the aforementioned Department; detailed information, including source and geographic origin for each genotype, is presented in Supplementary Table S1.

Genomic DNA from the 44 genotypes was isolated from young tomato leaves using the DNeasy Plant Mini Kit (Qiagen, Valencia, CA, USA). DNA quantity and purity/degradation was firstly checked on 1% agarose gel. In order to fit the standard parameters for Sequence Capture analyses, DNA concentration and quality were determined by Nanodrop spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA) and Qubit fluorometer (Life Technologies, Darmstadt, Germany) according to the manufacturer's requirements.

An inventory of 378 candidate genes representing possible targets for antioxidant metabolism in tomatoes was compiled from published literature, previous research^{31,32} and exploration of metabolic pathway databases (LycoCyc at http://solgenomics.net/and KEGG at http://www. genome.jp/kegg/). Genes were also selected to represent different gene family sizes. In total 235 out of the 378 genes completely represent 25 gene families. In particular, 171 genes belong to eight large gene families (we declared a big family if composed by more than 10 copies in the genome), 40 belong to seven medium gene families (between nine and four gene copies), and 24 belong to 10 small gene families (two or three copies). Six out of the 378 genes represent single copy genes. Moreover, in order to take into account variations in the regulatory region of each selected gene, a 3 Kbp promoter region was also included in the analysis. For genes with an intergenic distance shorter than 3 Kbp, only the related shorter portion of the promoter was considered. Supplementary Table S2 shows detailed information of the genomic regions considered.

2.2. Sequence capture design and sequencing

Probe design and gene enrichment were performed following the protocol provided with the solution-based Roche NimbleGen SeqCap EZ Library (Roche-NimbleGen, Madison, WI, USA). *Loci* coordinates of the genic and promotor regions were identified and submitted to Roche Diagnostics for probe design using NimbleDesign and SignalMap software (http://www.nimblegen.com/products/software/index.html). This probe set contains probes with up to 20 close matches in the genome as determined by the Sequence Search and Alignment by Hashing Algorithm (http://www.sanger.ac.uk/resources/software/ssaha/). Acco rding to NimbleGen specification, we considered a probe to match the genome if the variation ratio was <0.05. Following the SeqCapEZ protocol, 44 paired-end libraries were prepared using the Illumina Kapa Library Prep Kit and were multiplexed and sequenced with HiSeq 1,500 (read size:100 bp) according to Illumina specifications.

2.3. SUPER-CAP usage and procedures

SUPER-CAP is a bioinformatics tool specifically designed for accurate mapping, variant calling and reconstruction of private sequences using the variations detected from Sequence Capture data. SUPER-CAP includes an updated version of SUPER-W (release 4).²⁸ SUPER-CAP is a user-friendly web tool which only needs two input files to work: the captured region file in BED format and the filtered NGS reads in FASTQ format. The main steps and procedure underlying the SUPER-CAP tool, including the calling variants, the variants filtering and the targeted sequence reconstruction are presented in Figure 1.

SUPER-W has been modified to specifically handle sequence capture data. In this new version, SUPER-W uses as input the BED file of capture probe design (it can also handle a whole exome) and the filtered NGS reads. The first step is to map all the samples against a reference genome (specified by the user) with BWA (version 0.7.5; options used: mem:BWA-MEM algorithm).³³ The mapped files are processed for PCR duplicates (Picard, MarkDuplicates tool, version 1.118), filtered for quality (minimum quality required: 30), sorted and indexed.³⁴ The resulting BAM file is then used for the variant calling step. Small variations (SNPs and short deletion and insertion polymorphisms), large variations (deletions, inversions and duplications), or both, can be set for detection. The small variations are called with SAMtools³⁴ through an accurate and sensitive double calling step, while the structural variants are detected using LUMPY (version 0.0.11; options used: -mv 4 –tt 0 –pe –sr).³⁵

Once the variant calling step has been successfully completed, statistics on sequence capture experiment are calculated. The statistics report can be considered as a checkpoint, allowing the user to check the IN/OFF target read count, which highlights the specificity or sensitivity of the sequence capture experiment.

Raw genomic variants annotated by SUPER-W are then classified and filtered out. As default, only variants with a coverage $>6\times$ and a Phred quality >30 were used for subsequent analysis. Furthermore, variants overlapping each other are removed from downstream analysis as they cause a low confidence region. Moreover, a special option has been released to allow calling variants according to customized allele frequency (AF). As default value, variants with an AF between 0 and 0.2 are considered Ho for the reference allele, variants with a AF between 0.4 and 0.6 are considered He while variants with an AF between 0.8 and 1 are considered Ho for the alternative allele. Variants with an AF ranging out of boundaries set (i.e. between 0.2 and 0.4 and between 0.6 and 0.8) are called with an alert comment in the output VCF file. Coverage and quality filters as well as the AF parameters can be manually modified by the user. Finally, the filtered variants were classified according to type (SNP or



Figure 1. The SUPER-CAP tool. The main steps and procedures underlying the tool are graphically reported in the workflow.

INDEL). For SNP variants the breakdown into transitions and transversions was also determined. To improve the variant features, SnpSift and SnpEff³⁶ were implemented in the workflow, which allows imputing the gene region in which the variants fall (promoter, intron, exon, Untranslated regions (UTRs)) and the putative impact of the variants on protein functionality.

The high-quality variants, the reference genome and the BED file of the captured regions were used to perform a targeted sequence reconstruction. Using bcftools³⁷ and BEDtools³⁸ utilities, a new consensus sequence was created by applying the filtered variants to the reference sequence. He SNPs were introduced using IUPAC ambiguity codes and He INDELs were discarded. Through a liftover process, which recalculates the gene coordinates after the insertion or deletion of INDELs, a FASTA file with reconstructed sequences was created. Variants reconstructed were also displayed in the SUPER-CAP's browser results page, allowing the user to explore the type and the number of variants easily.

2.4. Web tool development and design

With the aim of supplying SUPER-CAP with a graphical user interface, we developed a NodeJS-based web tool (version 4.4.1. Available at https://nodejs.org/), available at 'http://supercap.sequen tiabiotech.com/'. Using this tool, the user can upload the two required files (the BED file of the target regions and the FASTQ files of the sequenced reads) and choose the analysis parameters (i.e. the quality thresholds for the filtering procedure and the AF range for the He /Ho assignation). An ID is given after completion of the procedure, which is required for accessing the results page. Both the reconstructed sequences in FASTA format and the whole variants in VCF format can be downloaded from the 'Download section'. In addition, the visualization of the reconstructed sequence is also shown in an interactive graphical interface in the results page. Here the user can explore, for each gene in each genotype, the type and the number of variants that have been incorporated and the effect they produced on the reference protein. General mapping statistics, including IN/ OFF targets, as well as the sensitivity of the experiment, are displayed in the same page.

2.5. Use of SUPER-CAP on tomato experimental data

The reads obtained by sequencing the tomato DNA samples were quality checked using FastQC (http://www.bioinformatics.babra ham.ac.uk/projects/fastqc/). Then, reads were filtered for quality and trimmed with the Trimmomatic tool (version 0.30; options used: windowsSize 3 -requiredQuality 22 -MINLEN 30).³⁹ By using the filtered reads, the following procedures were undertaken: reads falling in the designed regions with a PHRED quality >30 and a minimum length of 30 bp were mapped onto the tomato genome SL2. 50.40 Then, PCR duplicates and multiple mapped reads were removed by filtering on a meaningful MAPQ score of 30. Once the variants were called, in order to reduce the number of false positives calls, an additional filter was applied and only variants with coverage higher than $6 \times$ and a PHRED quality >30 were selected. The total numbers of He and Ho variants, as well as their ratio, were calculated with default parameters (0.4-0.6 as optimal AF range for the He calling). A Neighbour-joining dendrogram was built in TASSEL v.5⁴¹ by using the detected filtered variants, and the related tree was graphically refined using the Fig-tree software (http://tree.bio.ed.ac. uk/software/figtree/). SNPs and INDELs were also structurally and functionally annotated with SnpEff v4.2³⁶ using the tomato iTAG2. 40 annotation.

2.6. Validation of variants

To test the reliability of the adopted procedures, a validation of the variants detected was achieved by exploring publicly available data for tomato. In particular, data from a previous genotyping experiment which detected variants for 7,720 SNPs through the SolCAP array,^{30,32} as well as data concerning the resequencing of 360 tomato genomes,⁴² were considered. Of the 44 genotypes used in this study, 37 were common to those used in the SolCAP experiment and 9 in the 360 genome project. The validation was performed by comparing the number and the concordance of shared variations among the common genotypes. Only Ho variations were used for the validation. Intersection of the sets was carried out by intersectBed tool³⁸ and the comparison analysis by using VCFtools.⁴³

2.7. CNV and PAV identification

The coverage and total number of reads were evaluated for each sample. Alignments were visualized using the IGV browser version 2.3.3 (http://www.broadinstitute.org/igv/). CNV was computed following a modified procedure of Schiessl *et al.* $(2014)^{22}$ using a normalized read coverage for each captured region. CNV in a given target region was assumed if the ratio of normalized coverage (genotype)/normalized coverage (all genotypes) was <0.5 or >1.5, respectively. PAV was assumed if the ratio was <0.05.

3. Results and discussion

3.1. Sequence capture experiment

We designed a custom capture probe experiment by using a customizable and cost-effective approach based on Roche-NimbleGen SeqCap EZ technology. This framework helped to set up an *in silico* approach (SUPER-CAP) with the aim of facilitating further sequence capture studies.

This study investigated the sequence variation of a group of 378 genes and related regulative regions. The total size of the sequenced region was 2,338,578 bp, of which 1,353,817 bp corresponded to genic regions and 984,761 bp to their related promoter regions. Capturing probes were developed by Roche NimbleGen to target the specific regions. The final design was non-redundant and covered about 92.6% of the targeted regions. The 7.4% of non-covered regions represents those that did not match the minimum parameters for the capture and in the majority of the cases (>95%) these were promoter regions. It has already been reported that the cover design slightly varies depending on different factors, including the commercial technology adopted.⁴⁴

After the enrichment of the targeted regions, each capture library was sequenced in one Illumina HiSeq lane. The number of raw reads generated per sample varied between 1,103,352 and 2,683,868 (Table 1). Sequenced reads were analyzed in order to remove lowquality regions. After quality trimming, between 71 and 93% of the reads were retained and then used as input for SUPER-CAP. Data were then statistically evaluated for: (i) alignment rates, the number of mapped reads on the total reads, (ii) specificity, the number of reads that map to the targeted sequence and (iii) sensitivity, the percentage of targeted bases covered by sequence reads. The reads from samples 21A, 1A and 64A showed the highest alignment rates (>98%). In contrast, the lowest alignment rate was observed in genotypes 85A, 79A and 38A (<78%). The alignment success was however independent of the total number of reads. Specificity of the capture was imputed considering the number of reads in the target interval on the number of mapped reads. It showed an average value

Table 1. Alignment statistics for the 44 individual captures (Sample ID)

Sample ID	Raw reads	Mapped reads	Mapped reads Q > 30	Mapped reads Q > 30 WD	Alignment rate (%)	On target reads (%)	On target reads + 200 bp (%)	Mean depth of coverage
1A	1,344,456	1,327,567	1,269,157	1,251,088	98.74	75.45	76.06	43.2
3A	1,314,574	1,293,498	1,234,679	1,213,125	98.4	75.07	75.65	42.42
5A	1,413,706	1,391,398	1,331,163	1,310,092	98.42	75.94	76.55	45.97
8A	1,376,510	1,342,280	1,279,940	1,262,285	97.51	74.93	75.51	43.94
14A	1,580,074	1,231,066	1,142,665	1,128,063	77.91	67.22	67.75	39.67
15A	1,686,492	1,340,263	1,274,719	1,262,193	79.47	75.66	76.27	44.92
20A	1,480,340	1,162,656	1,101,676	1,056,514	78.54	71.73	72.28	40.27
21A	1,304,914	1,289,290	1,228,450	1,211,447	98.8	74.62	75.21	41.77
26A	1,103,352	871,462	830,910	821,995	78.98	75.72	76.35	30.03
27A	1,469,212	1,436,034	1,365,188	1,344,283	97.74	74.73	75.3	47.02
28A	1,354,892	1,332,311	1,264,586	1,190,895	98.33	70.39	70.87	40.76
30A	1,397,308	1,379,189	1,319,320	1,300,299	98.7	75.76	76.43	45.43
32A	1,893,224	1,480,474	1,410,253	1,394,575	78.2	74.55	75.18	48.69
34A	1.891.936	1,478,312	1,411,043	1.396.990	78.14	75.18	75.84	48.54
35A	1,788,530	1,398,636	1.333.126	1.316.721	78.2	74.51	75.16	46.02
38A	1.886.966	1.468.035	1.399.866	1.386.113	77.8	74.57	75.21	48.23
40A	2.651.822	2.575.073	2,463,126	2,432,466	97.11	76.67	77.25	88.92
41A	2.683.868	2.594.090	2,482,444	2.451.378	96.65	76.75	77.31	90.36
42.A	1.450.568	1.424.424	1.348.207	1.329.068	98.2	74.28	74.84	46.26
43A	2.082.552	2.006.847	1.921.991	1.899.495	96.36	77.02	77.59	70.57
45A	2,409,462	2.316.282	2.221.361	2,193,201	96.13	76.59	77.18	80.52
57A	1.297.772	1.275.552	1,216,245	1,198,785	98.29	75	75.56	42.68
64A	1 446 368	1 429 441	1 366 044	1 346 218	98.83	75 98	76.56	47.44
66A	2 267 108	2 176 890	2 081 441	2 057 807	96.02	76.59	77.17	74.86
70A	1 372 572	1 297 126	1 223 744	1 206 911	94.5	70.35	73.28	41 49
75A	1 799 384	1 396 835	1 323 958	1 312 523	77.63	73.81	74.38	45 59
78A	2 060 104	1,992,077	1 903 906	1,876,376	96.7	76.81	77.4	72 14
79A	1 666 724	1 295 896	1 235 321	1 224 643	77.75	75.02	75.63	44 71
85A	1,879,204	1,225,020	1 364 666	1 344 083	76.59	73.83	74 42	47.2
874	2 342 600	2 247 877	2 1 50 105	2 124 074	95.96	76.24	76.83	77.91
92 Δ	2,342,000	2,247,877	2,150,105	2,124,074	96.34	75.25	75.84	73.65
934	1 877 134	1 465 142	1 394 154	1 379 154	78.05	73.25	74.95	48 29
94 A	2 105 320	2 019 814	1 927 953	1 904 794	95.94	75.59	76.25	68 51
97A	2,105,520	2,012,014	2 299 398	2 270 879	96.51	76.11	76.72	82 59
99 1	2,407,574	2,402,528	2,277,376	2,270,079	96.01	76.11	77.11	81.97
1024	1 760 260	1 374 039	1 305 068	1 293 203	78.06	73.31	73.91	44 39
102/	1,700,200	1,974,037	1,305,008	1,275,205	95.77	71.5	73.21	61.9
105A	1,000,444	1,000,170	1,780,373	1,747,505	79.07	76.2	76.9	51.22
105A 109A	1,942,196	1,316,300	1,430,772	1,436,620	78.07	76.5	75.20	49.29
1111	2 5/1 20/	1,720,227	1 912 540	1,073,073	78.02	76 22	75.37	T7.20
1117	2,341,004	2,000,120	1,713,340	1,073,323	70.73	70.22	/0.82 71.2	00.21
113A 117A	1,04/,104	1,227,010	1,100,472	1,077,575	/ 7.47	70.75	/1.3	41.3
11/A 110A	1,682,246	1,333,089	1,2/3,3/3	1,262,226	/ 9.24	/6.42	//.03	44.83
1204	1,003,304	1,320,667	1,200,016	1,248,760	/7.3	/3.09	/3./2	43.63
12UA	1,031,764	1,308,823	1,247,77	1,237,434	/7.23	70.03	/ 0.64	43.62
Mean	1,806,624	1,602,892	1,52/,880	1,303,873	88./2	/4.80	/3.40	55.89

The number of raw reads (Raw reads), mapped reads (Mapped reads), mapped reads with quality Q > 30 (Mapped reads Q>30) and without duplications (Mapped reads Q > 30 WD) were reported. For each sample, the alignment rate (Alignment rate), the specificity (On target reads), the specificity including the flanking regions of 200 bp (On target reads + 200 bp) and the average depth of coverage (Mean depth of coverage) are also reported.

of 74.80%. The minimum value 67.2% was detected for the genotype 14A and the maximum (77.02%) for the genotype 43A. In addition, considering an extension of flanking regions of 200 bp, the specificity was found to vary only slightly, showing an average increase of <1%. The range, considering flanking regions, was indeed from 67.75 to 77.59%. variant-error rate and the assembly gaps, and to obtain the correct call of variants, in particular for the He ones.⁴⁵ It has indeed been demonstrated that for short Illumina reads a coverage comprised between $30 \times$ and $55 \times$ appears necessary to correctly identify SNVs and small INDELs with a proper degree of reliability.^{45,46}

Even though investigators are seeking experimental designs that generate robust scientific findings at the lowest sequencing cost, an adequate depth of coverage is very important in order to reduce the In our experiment, the normalized mean coverage of the total targeted regions showed an average value of 53.89×, allowing very accurate detection of variants. Analysing the sensitivity of the experiment (Supplementary Fig. S1), which shows how well the

targeted region is covered for reads at depths from $1 \times$ to the max depth (>150×), we observed that at a minimum coverage of $1 \times$ about 95% of the designed target regions resulted completely covered. Between 91 and 94% of the target was covered when a minimum coverage of ten reads was applied. At $20\times$, 76–93% of the target regions was still covered.

Although Sequence Capture experiments represent a promising technology, only few studies have used Sequence Capture in plants (<20 in PubMed in April 2016). Probably the lacking of straightforward procedures for data analysis makes this technology still emerging. For this reason, SUPER-CAP, representing a user-friendly tool, could boost the use of this technology also for plant organisms. Moreover, in this study, the high rates reached for the capture parameters, in particular for the specificity (>74%), highlighted the high efficiency reached by combing an accurate probe design with a high performant aligner included in the pipeline (BWA), for which high confidence of sequence alignments has been already reported.¹⁹ This makes the performances obtained in this study in terms of alignment rate and sensitivity in line with performances reported for other organisms, for example humans.44

3.2. Variant calling and reconstruction of private sequences

SUPER-CAP allowed to detect 25,654 unfiltered variants in the targeted regions considered. Then the filtering procedure allowed to

reduce the number of false positives/low-quality variants. Only variants exhibiting coverage higher than 6× and PHRED quality higher than 30 were selected. Even if a different threshold could be applied to the filtering procedure, we observed that in our experiment a depth coverage of six represented the minimum coverage to maximize the PHRED quality (Supplementary Fig. S2). Subsequent criteria were adopted to appropriately detect the number of He variants. A preliminary survey was carried out in order to estimate the number of He variants in accordance with the range of the AF (AF of mapped reads for each locus) (Supplementary Fig. S3). As expected, an inverse correlation between the range of AF and number of He variants was observed. This indicated that the set-up of the AF-range deeply affects the estimation of the He variants in the calling procedure. In order to make this sensitive phase of the variant calling easily customizable, we allow the SUPER-CAP user to set their own AF range. In this study, we set an AF range of 0.4-0.6 for the He variants, in line with a previous study in tomato.¹⁰ At this threshold, 86,120 variants in 14,116 sites were scored. Only 5% (4,970) of the total variants in 2,543 sites resulted to be He. Looking at the distribution of the He variants, we observed that they were not uniformly distributed across the chromosomes and genotypes. Chromosome 11 harboured the highest number of He variants per Mbp of captured region (Supplementary Fig. S4) while chromosomes 10 and 0 harboured the lowest number. Considering the 44 samples analysed, the heterozygosity ranged from 1.1 to 47% (Fig. 2), with a minimum value for sample 99A and a maximum value for sample 85A. Sample



Figure 2. Distribution of variants detected for each genotype according to the type of variants. Proportion of Ho and He SNP and INDEL are reported as well as the number of private variants. A dendrogram built by using the complete set of variants shows the relationship among the genotypes.

85A showed an unexpected high percentage of residual heterozygosity, suggesting a possible sampling error of the plant material for this genotype (which could be mixed with other genotypes or to a residual segregation during the germoplasm conservation). For this reason, the sample was excluded from subsequent analyses.

Figure 2 shows the number of variants detected for each sample, with respect to the proportion of SNPs and INDELs at both Ho and He level. Although INDELs are present at lower rates than SNPs, small INDELs represent functionally important types of genomic variation.^{47,48} It is reported that, compared with other sequencing technologies, Illumina does not appear to have high error rates in homopolymer regions.⁴⁹ However, in this study the INDELs surveyed were obtained after discarding positions overlapping with homopolymers >7 bp, both in the tomato reference and in resequencing sequences. This allowed to avoid possible errors due to technical bias and enabled to collect a final high-quality set of variants.

As a whole, the sample with the highest number of variants was 42A, which harbours >4,000 SNPs and about 1,000 INDELs. The sample with the lowest number of variants was 102A which exhibits 1,123 variants between SNPs and INDELs. On average, 1,357 SNPs *per* sample were scored, of which 6.8% turned out to be He. The average of INDELs *per* sample turned out to be about 600, and only 3% were He.

The high depth of coverage reached in this experiment supported the detection of rare variants. It has been suggested that rare or low-frequency variants, which are not fully captured using other conventional genotyping technology, could largely contribute to explain the effects of some traits⁵⁰ or to characterize their genetic architecture.⁵¹ In this study 5,441 Ho and 442 He genotype-specific variants were identified. Among these, 1,965 Ho and 216 He variants resulted still private variants when compared with external public repository for SNPs in tomato (360 tomato genomes⁴²). This could be of interest when designing specific SNP markers or for studying the potential functionality of specific haplotypes in the landraces collection used in this survey. However, the rare variants identified alone do not seem to explain the phylogenetic differences observed (Fig. 2). The genetic divergence was associated with the high number of private alleles only in a few samples (i.e. 103A and 97A).

The possibility of reconstructing private sequences from highthroughput resequencing data is still a challenge today.²⁸ For this reason, the last step in the SUPER-CAP workflow allows to specifically insert the detected variants into the sequence of each sample. This step, which integrates the single information of the variants in a genotype-specific gene-based view, enables to obtain a necessary starting point for further functional validation analyses. In this study, a total of 16,254 regions (378*43), each including the gene and the related promoter region, were reconstructed. In order to easily exploit the combined effect of multiple variants in the same protein, reconstruction of the CDS was also performed. Future works, benefitting from this effort, will evaluate the combined effect of the variants for each gene in order to find relevant association with antioxidant content in tomato.

3.3. Validation of variants

In order to validate the variants detected, we performed a survey by exploring the available resources already present for tomato. About 8,600 variant sites detected in the present study overlapped with those reported in the genome resequencing data of 360 tomatoes.⁴² Most of the non-overlapping variants were excluded since they classified as rare variants in the experiment, being present only in specific

samples. Moreover, by using nine common accessions, an analysis on the concordance of the variant calling was performed. Each of the nine samples shared a different number of variants, ranging from 191 to 985. Taking into account the Ho alternative variants only, an average of 95% concordance was found. Another validation was carried out by comparing SNPs detected on a tomato population genotyped using a SolCAP arrays,^{30,32} which included 7,720 SNPs. Only 170 of the 7,720 SNPs fall in the targeted region considered in the present study, since the targeted regions represent ~1% (378/ 34,725) of the tomato genes. Of these 170 SNPs, a subset of 97 belongs to common accessions and was used for validation. On average, a concordance >95% was observed. In most cases the discrepancies were genotyped as He either on the SolCAP array (2.5%) or in the Sequence capture (0.6%). When excluding the He cases, the similarity increased to 98%.

3.4. CNV and PAV

CNV and PAV have been recently reported as sources of important phenotypic variation in plants.⁵²⁻⁵⁴ The adequate depth coverage reached in this study allowed to hypothesize on the possible structural variations occurring among the samples considered. To do this, CNV and PAV analyses were performed to detect additional or deleted homologous loci of the 378 genes among the 43 samples, by analysing the normalized depth coverage of each sample. The comparison of read depths along the loci revealed at least 10 loci with a significant variation in depth in at least one line. In particular, nine genes (four Pectinesterases, two Phenylalanine ammonia-lyase, one Reductase, one Inositol-3-phosphate synthase, one Caffeoyl-CoA Omethyltransferase) showed a CNV, and one gene (Dehydroascorbate reductase) showed a PAV (Supplementary Table S3). Three genes showed a putative reduction of the CNV (ratio <0.5) while six showed an increase (ratio >1.5). The very low number of reads captured for two samples (103A and 15A) for the gene Solyc09g056180 was associated with a PAV and we assumed that this gene was deleted in the two samples. In addition, an experimental validation by genomic PCR corroborated this result, highlighting that no amplification of the Solyc09g056180 was present for these two genotypes as showed in Supplementary Figure S5. The analysis of the coverage revealed a uniform high level of coverage across all the samples for seven genes (Solyc00g027770, Solyc00g282510, Solyc00g030510, Solyc02g075620, Solyc03g042560, Solyc03g036470, Solyc03g07 1860). Indeed, these seven genes showed levels of coverage ranging between 104.58× and 192.48×, representing an increase of two to four fold compared with the average coverage encountered among all the genes $(53.89 \times)$. This could be due to the lack of regions in the current release of tomato. A high number of reads collapsing on specific regions could be caused by unassembled/misassembled regions in the genome.²⁹ To validate this unexpected high coverage, we evaluated the depth of coverage of these genes in two further independent resequencing experiments (Heinz and Moneymaker) by using publicly available data (variation data from SGN databases). We observed that significant high coverage was obtained for three genes in at least one of the two resequencing experiments and one gene in both the resequencing experiments (data not shown). This corroborates that the reference genome could carry misleading sequences. An additional indication of this hypothesis, as also discussed below, come from the high ratio of He observed for these genes, showing that similar but not identical reads were mapped on the reference regions.

3.5. Heterozygous variants into gene families

He variant assignments represent one crucial difficulty in sequence experiments.^{10,29,55} Since one of the causes of spurious He call was hypothesized to derive from duplicated genes, a specific investigation was carried out, aided by the high quality of variants detected by SUPER-CAP. Previous studies reported that false-positive signals could arise in regions with low complexity,⁵⁶ or result from misalignment of multiple copies of genes, paralogues, or pseudogenes⁵⁷ but, to our knowledge, no study has analysed the behaviour of the He variants in the gene family context.

In our collection, 235 out 378 genes completely represent 25 gene families. In particular, 171 genes belong to eight large gene families, 40 belong to seven medium gene families, 24 belong to 10 small gene families and six are single copy genes. In order to estimate the number of He variants in each group we evaluated (Fig. 3) the average He density (number of He variants for Kbp) for each gene family. On average, a lower number of He variants was identified in the single genes compared with the gene family groups. Significant differences were observed at t-test between single genes and Large, Medium and Small gene families (P = 0.007) but not among gene families themselves (P > 0.05). Although these results evidence that gene families, despite the number of copies, are more prone to produce He variants compared with single genes, the high standard deviation detected showed that differences seem to be associated with specific cases in each gene family and not due to a general behaviour. It appears evident that in each gene family only specific genes showed a very high level of heterozygosity in almost all the genotypes (Supplementary Fig. S6). In particular it was evident that Solyc03g042560, Solyc00g027770 Solvc00g282510. and Solyc01g091060 (belonging to large gene families) and Solyc12g098090 (belonging to a medium gene family) showed a skewed rate of He variants. Some possible explanations of this result were hypothesized. In particular, exploring if neighbouring regions of these genes showed a similar rate of heterozygosity (by using information from the 360 genome project), we evidenced that a similar high rate of He variants was found only for Solyc03g042560 in a region spanning ~ 300 Kbp.

Another source of false positive He variants might be due to duplicated regions found in the samples that are not found in the reference genome. In such cases, the variants will be due to reads from the two copies (or more) that are piled in the only copy found in the reference genome. Such biases could occur because of the limited number of genotypes on which the original reference sequence was based, or sequencing and alignment errors (Lander *et al.*, 2001). Fortunately, these cases can be highlighted looking at excesses of depth of coverage. Three out of the five genes (Solyc00g282510, Solyc00g027770, Solyc01g091060) exhibiting a high He density showed a high depth of coverage in parallel (as also evidenced in the previous paragraph), generally doubled or tripled in comparison to the average value of all the samples. This proves that the tomato reference genome might carry misleading sequences for those regions and that the He calls observed are likely to be false-positives.

3.6. Variants classification and annotation

As an additional step, the functional annotation of the variants has also been investigated by using SnpEff and SNPsift programs. The SNPs discovered were classified according to their type (transition vs transversion and insertions vs deletion) and the genomic region they were found in. A significant number of the SNPs discovered (6,098, 44.2%) were located in the genic region and, among these, about 21% were found within exons, 73% within introns, 1.73% within the 5'-UTR and 3.68% within the 3'-UTR (Fig. 4). The remaining 7,674 (55.8%) were found to be within the promoter region. Considering the number of variants per Kbp, eight variants per Kbp were found in the promoter region on average, while only 2.2 variants per Kbp were found in the exonic regions. Introns and UTRs showed more than double the number of variants per Kbp compared with exonic regions. Genic and intergenic variants were classified as transition/transversion and insertion/deletion typologies. Transitions represent approximately 64.54% of all post-filtered SNPs (11,495), with 44.2% (5,160) of them being located in genic regions. Deletions in average were more represented than insertions, particularly in promoter and intron regions. Then, the 6,098 variants of the genic regions were annotated according to their putative effect on the protein by using SnpEff v4 (Table 2). The goal of annotating variants is to provide a prediction of which ones are functionally relevant. Our detected variants were assigned to a diverse range of functional classes, with the majority (78%) classified as 'modifier', therefore without predictable effects being located in the UTR regions or in the introns. These may have little or no effect on the phenotype. Among the remaining variants, about 12% showed a 'low effect', since they



Figure 3. Average He variants density (He density) for each gene family (reported as EC number) in each category (LARGE, MEDIUM, SMALL, SINGLE). He density is expressed as the average number of He variants for 10 Kbp. Bar errors represent SD of the mean.

1		PROMOTER	5'-UTR	EXON	INTRON	3'-UTR
	Total captured region (bp)	972,923	23,388	575,336	791,726	48,357
•	Average length per locus (b	p) 2,567	61	1,518	2,088	127
•	No Variant sites	7,674	104	1,309	4,460	225
•	No variants per Kbp	7.88	5.60	2.27	5.63	4.65
•	Variant Type:					
	- No SNP	6,335	67	1,255	3,658	180
	- Ts	4,148	37	808	2,315	112
	- Tv	2,151	30	447	1,343	68
	- No INDEL	1,339	37	54	802	45
	- INS	589	13	27	324	18
	- DEL	750	24	27	478	27

Figure 4. Distribution and classification of the discovered variants according to type (transition vs transversion and insertions vs deletion) and according to the genetic feature they were found in (promoter, exon, intron, UTRs).

 Table 2. Distribution of 6,098 genic variants per type of predicted

 effect (PREDICTED EFFECT) on the relative protein as predicted by

 SNPeff.

PREDICTED EFFECT	VARIANTS No			
Modifier	4,765			
3' UTR variant	256			
5' UTR variant	155			
Intron variant	4,354			
Low	729			
5' UTR premature start codon gain variant	13			
Splice region variant & intron variant	115			
Synonymous_variant	601			
Moderate	576			
Missense variant	559			
Inframe deletion	7			
Inframe insertion	10			
High	28			
Frameshift_variant	14			
Splice acceptor variant	2			
Stop_gained variant	9			
Stop_lost variant	3			
Total	6,098			

were variants in coding regions that did not change the amino acid sequence (synonymous variant). About 9% showed a 'moderate' effect. These variants are predominantly non-synonymous amino acid changes (missense variant) and in few cases in-frame deletion/insertion. They are the most likely candidates for causal mutations, since they could alter the structure and function of relevant proteins. Last, about 1% of the variants were predicted to have a 'high effect'. In particular, 14 variants were predicted to cause a frameshift: two of them a splice site acceptor modification, and nine and three a start or stop codon gain or loss, respectively. On average, we observed that the percentage of INDELs with 'high effect' was higher than for SNPs, since an INDEL may rapidly cause a frameshift in the sequence. The list of the variant with a moderate or a high effect with respect to the gene they affected is provided in Supplementary Table S4.

Thus, a larger effort would be warranted to study potential links between the identified variants and trait variation in tomato, and to determine how they affect the regulation of biological pathways and processes. The variants selected or prioritized in this way would be highly preferred marker sets to be subjected to association studies using suitably larger populations, for which panels with considerable genotype and phenotype information has already been collected.^{30,32}

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Conflict of interest

None declared.

Supplementary data

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

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