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Timothy L Fitzgerald1*, Kemal Kazan1, Zhongyi Li2,3, Matthew K Morell2,3, John M Manners1

Abstract

Background: Mutational inactivation of plant genes is an essential tool in gene function studies. Plants with inactivated or deleted genes may also be exploited for crop improvement if such mutations/deletions produce a desirable agronomical and/or quality phenotype. However, the use of mutational gene inactivation/deletion has been impeded in polyploid plant species by genetic redundancy, as polyploids contain multiple copies of the same genes (homoeologous genes) encoded by each of the ancestral genomes. Similar to many other crop plants, bread wheat (Triticum aestivum L.) is polyploid; specifically allohexaploid possessing three progenitor genomes designated as ‘A’, ‘B’, and ‘D’. Recently modified TILLING protocols have been developed specifically for mutation detection in wheat. Whilst extremely powerful in detecting single nucleotide changes and small deletions, these methods are not suitable for detecting whole gene deletions. Therefore, high-throughput methods for screening of candidate homoeologous gene deletions are needed for application to wheat populations generated by the use of certain mutagenic agents (e.g. heavy ion irradiation) that frequently generate whole-gene deletions.

Results: To facilitate the screening for specific homoeologous gene deletions in hexaploid wheat, we have developed a TaqMan qPCR-based method that allows high-throughput detection of deletions in homoeologous copies of any gene of interest, provided that sufficient polymorphism (as little as a single nucleotide difference) amongst homoeologues exists for specific probe design. We used this method to identify deletions of individual TaPFT1 homoeologues, a wheat orthologue of the disease susceptibility and flowering regulatory gene PFT1 in Arabidopsis. This method was applied to wheat nullisomic-tetrasomic lines as well as other chromosomal deletion lines to locate the TaPFT1 gene to the long arm of chromosome 5. By screening of individual DNA samples from 4500 M2 mutant wheat lines generated by heavy ion irradiation, we detected multiple mutants with deletions of each TaPFT1 homoeologue, and confirmed these deletions using a CAPS method. We have subsequently designed, optimized, and applied this method for the screening of homoeologous deletions of three additional wheat genes putatively involved in plant disease resistance.

Conclusions: We have developed a method for automated, high-throughput screening to identify deletions of individual homoeologues of a wheat gene. This method is also potentially applicable to other polyploid plants.

Background

Identification of plant gene function is important not only to better understand how plants grow and develop, but also for potential exploitation of this information for molecular and/or traditional crop breeding. However, extensive gene duplication and polyploidy events during evolution mean that plants carry multiple copies of the same gene as well as large numbers of closely-related genes [1]. Therefore, for gene function studies, it is important to determine whether different copies of the same or related genes are functionally redundant or whether polymorphisms have been selected through evolution and led to functional specialisation [2,3]. Reverse genetics approaches to answering these questions range from the development of transgenic plants where individual gene copies are silenced [4,5] to the identification of mutations where gene copies have been either modified, inactivated by sequence alterations [6-8], or completely deleted [9]. For the model plant Arabidopsis, the availability of large numbers of T-DNA (or transposon) insertion lines makes it increasingly
likely that knock-out insertion mutants can be identified for specific copies of selected closely-related genes to test for redundancy or a specific function [10]. In many other plants, including most of the important crop species, this task is not so straightforward and new technical approaches for the identification of mutations in specific gene copies are needed [11].

Mutant populations of many crop species including wheat have been developed through chemical mutagenesis, and mutations in a range of target genes have been identified in such populations using ‘Targeting Induced Local Lesions IN Genomes’ (TILLING) techniques. In TILLING, a gene region is amplified from pools of genomic DNA, amplified fragments allowed to re-anneal, and mutations detected by testing for mismatches between wild type and mutant heteroduplexes [13,14]. This technique is capable of detecting the point mutations, re-arrangements, and small deletions [6] that chemical mutagens predominantly produce [13], but it is not directly applicable to screening for whole-gene deletions. An additional technical difficulty in applying TILLING to polyploid crop species such as wheat is that TILLING requires independent amplification of homoeologous gene copies to prevent heteroduplexes forming between these closely-related sequences, as this can confound the subsequent data analysis. Therefore, the development of new high-throughput methods that allow simultaneous detection of mutations in homoeologous genes in a single assay, as has recently been reported [15], is certainly highly desirable for functional genomics studies in wheat and other polyploid crop species.

Recently, physical mutagenesis such as heavy ion irradiation (HII) has been used to generate mutant populations of model, crop and ornamental plant species [16-18]. In contrast to chemical mutagenesis (e.g. ethyl methanesulphonate EMS treatment) which generates frequent point mutations and small deletions, HII generates a relatively high frequency of large (whole gene) deletion mutations. However, the difficulty of detecting mutants that contain deletions of homoeologues of target genes has been a major factor limiting the application of this method to polyploid species.

Bread wheat (Triticum aestivum L.) is one of the most important food crops in the world and is hexaploid, comprising the A, B and D genomes with theoretically all important food crops in the world and is hexaploid, comprising the A, B and D genomes with theoretically all homoeologues in wheat [19]. Similar to other plants, availability of genetic mutants that facilitate understanding of gene function is highly desirable in wheat. In addition, new genetic variation in mutants can be exploited in traditional plant breeding [6,11]. In wheat, well-recognised examples of the types of genes that are candidates for mutational inactivation are those that encode allergenic and nutritionally poorly tolerated grain proteins [19] as well as grain quality genes such as those that influence starch composition [20].

An emerging new category of genes for which null mutations will be sought are disease susceptibility genes [21-23]. It has become apparent that pathogens often exploit essential cellular processes of their hosts for their own benefit during infection and that multiple host genes may be required to permit a susceptible infection [22]. We have recently demonstrated that inactivation of the PFT1 gene of Arabidopsis led to increased resistance to the fungal pathogen Fusarium oxysporum [23]. The PFT1 gene was initially identified as a positive regulator of shade avoidance [24], and subsequently shown to encode a subunit of the plant Mediator complex - a conserved multi-protein complex involved in the fine-tuning of gene expression in all eukaryotes [25]. We are interested to test whether the inactivation of orthologues of PFT1 in wheat would have a similar effect on fungal disease resistance. The PFT1 gene is highly suited for reverse genetic analysis and the development of a new screening method, because it is a single copy gene in the genomes of diploid species such as Arabidopsis and rice, and a full-length cDNA sequence from wheat is available [23].

In the present study, we have developed a TaqMan single nucleotide polymorphism (SNP) detection method suitable for high-throughput screening of a HII population of hexaploid wheat to identify homoeologous deletion mutants. This method relies upon uniquely fluoro-labelled homoeologue-specific TaqMan SNP-detection probes to identify the presence or absence of homoeologous copies of a gene of interest. Although the TaqMan SNP-detection PCR system has been widely used as a high-throughput method for allelic discrimination in humans [26,27], this method has not been used extensively in plants, with some limited application to the differentiation of SNPs in genes that confer herbicide tolerance [28,29]. The application of TaqMan probes to screening for gene deletions in Drosophila has been demonstrated [30]. However, the methodology used in that study differed significantly to the method presented here as gene specific primer/probe combinations with generic probe fluoro-labelling were used in Drosophila to assess the presence of diploid loci by comparing fluorescence from individual gene specific qPCRs on the same DNA samples.

Here we describe in detail the design, optimization, validation, and application of a new method for the identification of deletions for each of the three homoeologues of a wheat gene, using TaPFT1 as a model. Using this method we have subsequently identified multiple deletion mutants for three additional wheat genes putatively involved in plant defence.

Results
Identification of TaPFT1 homoeologues in wheat
Based on an existing full-length cDNA sequence for a wheat orthologue (UniGene Ta.39294) [16] of the
Arabidopsis PFT1 gene in conjunction with the conserved intron-exon structure that exists in the orthologous single copy rice gene for PFT1 (Os09g13610), it was possible to design PCR primers (see Materials and Methods) that would generically amplify overlapping fragments of all homoeologous copies of TaPFT1. Multiple (30-60) DNA fragments amplified with conserved TaPFT1 primers from the wheat cultivar Chara were cloned and sequenced. This resulted in the identification of five unique contigs (designated TaPFT1a - TaPFT1e; see Methods) putatively corresponding to sequences that partially spanned between exon 4 and exon 15 of the 15 predicted exons of three TaPFT1 homoeologues (Additional file 1). The consensus exon-intron structure of the TaPFT1 gene in regions used for the design of the TaqMan probe assay and a Cleaved Amplified Polymorphic Sequence (CAPS) assay for validation are depicted diagrammatically in Figure 1. Sequences corresponding to TaPFT1 homoeologues in Chara were deposited into the NCBI database (see Methods for accession numbers). The total polymorphism amongst TaPFT1 homoeologous sequences is given in Table 1.

Probe-Based Detection of TaPFT1 Homoeologues

The high-throughput technique we describe here for reliable amplification of TaPFT1 homoeologues from wheat is modified from a dual-labelled probe SNP detection assay originally reported by Livak [26]. Based on sequences obtained for putative TaPFT1 homoeologues, a region of TaPFT1 suitable for the design of short fluorescent hybridisation probes that would distinguish different TaPFT1 homoeologues was identified (Figure 1) and three homoeologue specific TaqMan probes labelled with the fluorescent dyes FAM, NED and VIC were designed and synthesised. These probes were then used in qPCR together with a pair of conserved flanking PCR primers that would simultaneously amplify all three TaPFT1 homoeologues (Figure 2A). As shown in Figure 3, this assay successfully amplified TaPFT1 homoeologues from the genomic DNA isolated from Chara.

Validation of Assay

Before an extensive screening of mutagenized wheat populations was undertaken for identification of TaPFT1 deletion mutants, we hoped to validate the specificity of this assay by determining the chromosomal location of TaPFT1 using genetically well-characterized wheat lines (data not shown). We obtained a series of nullisomic-tetrasomic wheat lines, each missing a single chromosome (nullisomic) substituted by an additional copy of a homoeologous chromosome (tetrasomic) [31,32] as well as a comprehensive smaller deletion series of wheat lines [33]. These lines were generated using the cultivar ‘Chinese Spring’. The probe assay functioned successfully against wild type ‘Chinese Spring’ (Figure 4A), indicating that sequence in the targeted region was conserved between ‘Chara’ and ‘Chinese Spring’. This was confirmed in silico by performing a BLAST search of 5× Chinese Spring Whole Genome Sequence 454 raw reads (available at http://www.cerealsdb.uk.net) with TaPFT1 sequence obtained from Chara - reads with identical sequence to each of the homoeologous sequences in the region of assay design were detected. As shown in Figure 4, wheat lines individually missing chromosomes 5A, 5B and 5D lacked fluorescence from VIC-, FAM- and NED-labelled probes, respectively. This indicated that TaPFT1 was located on chromosome 5. The location of TaPFT1 on homoeologous chromosomes 5A, 5B, and 5D, respectively, was confirmed independently using the TaPFT1 CAPS screen (Figure 5A and 5B), which was also designed based on the obtained ‘Chara’ sequence (see Materials and Methods), but shown to function successfully for ‘Chinese Spring’ samples, as confirmed by BLAST searches at http://www.cerealsdb.uk.net.

To more specifically determine the genomic location of TaPFT1 on chromosome 5, we screened additional
deletion lines that contain smaller deletions on homoeologous chromosome 5A [33]. These experiments revealed that one line, 5AL-14, lacked the A genome homoeologue of \(\text{TaPFT1}\) designated here as \(\text{TaPFT1-A}\). 5AL-14 was the largest long-arm chromosome 5 deletion available in the series and the absence of the amplification of \(\text{TaPFT1-A}\) indicated that this gene was located between the break point of 5AL-14 (FL: 0.11) and the break point of the next largest chromosome 5 deletion, 5AL-12 (FL: 0.35) [33]. Together, these experiments also confirmed that the assay was able to reliably detect presence and/or absence of \(\text{TaPFT1}\) homoeologues in wheat.

**Identification of \(\text{TaPFT1}\) deletion mutants**

We next applied this assay for the identification of \(\text{HII}\) mutants that lack at least one of the \(\text{TaPFT1}\) homoeologues. From the screening of approximately 4500 individual \(\text{HII}\) mutant (M2 generation) lines of Chara, nine ‘definite’ mutants were detected, with an additional two ‘possible’ mutants identified. Mutants were considered ‘definite’ if efficient fluorescence increases for two of the three probes, with \(C_T\) (cycle thresholds) below 30 was observed, in conjunction with very low or no fluorescence increase for the third probe for both replicates. Mutants were considered ‘possible’ if one replicate failed, or if fluorescence from probes had a \(C_T\) value greater than 35, indicative of very low DNA concentration in the sample. Of the screened mutants, five samples (BW18M3-444, BW19M3-132, BW20M3-616, BW22M3-85, and BW22M3-127) lacked fluorescence from VIC-labelled probe 2, indicating the absence of \(\text{TaPFT1-A}\) (Figure 6A); two samples (BW18M3-157 and BW19M3-146) lacked fluorescence from FAM-labelled probe 1, indicating the absence of \(\text{TaPFT1-B}\) (Figure 6B); and four

| Table 1 Sequence diversity identified amongst the three homoeologues of \(\text{TaPFT1}\) in the sequenced regions |
|---------------------------------------------------------------|
|                    | base pairs | SNPs | indels | all polymorphism | SNPs/100 bp | indels/100 bp | all poly/100 bp |
|---------------------|------------|------|--------|------------------|-------------|--------------|----------------|
| exon sequence       | 1126       | 21   | 8      | 29               | 1.9         | 0.7          | 2.6            |
| intron sequence     | 4368       | 214  | 51     | 265              | 4.9         | 1.2          | 6.1            |
| total sequence      | 5494       | 235  | 59     | 294              | 4.2         | 1.1          | 5.4            |

**Figure 2** Alignment of \(\text{TaPFT1}\) homoeologues. ‘A’, ‘B’ and ‘D’ \(\text{TaPFT1}\) homoeologue sequences in regions used for design of the probe assay (A) and the CAPS assay (B), respectively (see Figure 1) were aligned. Forward and reverse primers used in the probe assay are highlighted in blue. Polymorphisms are highlighted in red. In (A), homoeologue specific probe sequences and in (B), restriction enzyme (Hpy166I) cleavage sites are highlighted in yellow.
samples (BW19M3-325, BW21M3-723, BW21M3-734 and BW19M3-757) lacked fluorescence from NED-labelled probe 3, indicating the absence of TaPFT1-D (Figure 6C).

Of the eleven putative mutants, seed stocks were not available for BW18M3-444 and BW19M3-132; and seeds of BW19M3-146 failed to produce viable seedlings. DNA extracted from the remaining lines were screened using the probe-based screen in conjunction with the TaPFT1 CAPS screen and in this case, CAPS gels were also blotted and hybridised to a TaPFT1 probe to ensure specificity (Figure 7). The results of CAPS screening and hybridisation (Figure 7) demonstrated that genome-specific bands present in Chara were missing in the designated mutants for samples even including the ones that were considered possible in the initial screening.

Application of method to identification of deletions in other target genes
To confirm that the method described here performs routinely, we have subsequently chosen three additional wheat genes putatively encoding negative regulators of plant disease resistance and screened our HII mutant population to identify homoeologue deletions in these genes. The genes selected were: TaNFXL1, TaCesA8, and TaPLD1. TaNFXL1 is the orthologue of the Arabidopsis AtNFXL1 gene demonstrated to be a negative regulator of salicylic acid (SA) accumulation and SA-dependent defence genes. Arabidopsis mutants lacking AtNFXL1 show increased resistance to the compatible pathogen Pseudomonas syringae pv. tomato strain DC3000 (Pst DC3000) [34]. TaCesA8 is the orthologue of the Arabidopsis CesA8 gene shown to encode a susceptibility factor for the bacterial pathogen Ralstonia solanacearum and the necrotrophic fungus Plectosphaerella cucumerina. Resistance in cesa8 mutants is believed to be conferred via an increase in ABA-dependent defence gene expression, and alteration of cell wall structure, leading to the activation of novel defence pathways [35]. TaPLD1 is the orthologue of OsPLD1, a rice phospholipase D gene demonstrated to act as a negative regulator of resistance to Magnaporthe grisea and Xanthomonas oryzae pv oryzae. Knockdown of OsPLD1 results in induction of a range of defence related genes in rice [36].

cDNA sequences corresponding to putative wheat orthologues of these additional target genes were identified using Arabidopsis and/or rice protein sequences as queries in tblastn searches of the NCBI wheat EST sequence database. Sequences for rice orthologues of AtNFXL1 and AtCesA8 were obtained by performing a tblastn search against the NCBI rice cDNA sequence database. Full genomic sequences are available for all annotated rice cDNAs. Therefore, putative intron/exon structure of corresponding wheat genes was determined by aligning wheat cDNA sequences to rice genomic sequences for all genes. Based on these alignments PCR primers were designed. PCR products were cloned and more than 30 individual clones were sequenced for each gene and, based on these sequences, the three putative homoeologous sequences were deduced. Using these homoeologous sequences, assays were designed as for TaPFT1 and applied to the screening of the HII wheat lines. Consequently, 14, 19 and 11 deletion mutants were identified for TaNFXL1, TaCesA8, and TaPLD1, respectively, demonstrating that the assay is broadly...
applicable to deletion screening in wheat. Details of the design and application of these assays are provided in Table 2.

Discussion

A greater understanding of the genetics underlying the phenotypic properties of crop species has improved our ability to manipulate genes responsible for unwanted characteristics. Genes encoding certain grain proteins which can be allergenic or nutritionally poorly tolerated are well recognised examples of genes with unwanted characteristics [19]. Many desirable characteristics are also known to be controlled by recessively inherited loci [37-39] some of which have been shown to be non-functional alleles [40,41]. Furthermore, in the last decade ‘negative regulators’ of plant resistance have been reported. Loss-of-function mutants of these genes show increased resistance to specific pathogens [21-23]. It therefore seems likely that reverse genetics approaches that exploit the use of gene deletions in crop improvement will increase, especially whilst GM approaches for specific gene inhibition may be restricted in commercial use due to prohibitive legislation in many parts of the world.

In this article we outline a TaqMan SNP-detection probe-based assay optimized for high-throughput screening to identify whole-gene deletion mutants in wheat and other polyploid species. The following strategy was followed to develop and validate the assay. First, sequence analysis was undertaken on the TaPFT1 gene to identify homoeologous sequences. This sequence information was then used to design a qPCR assay using specific TaqMan SNP-detection probes that would discriminate the three TaPFT1 homoeologues in a single reaction. To confirm the ability of the assay to detect wheat lines missing a specific TaPFT1 homoeologue, it was applied to the screening of nullisomic-tetrasomic wheat lines [31,32]. Chromosomal locations of TaPFT1 homoeologues were thereby identified, and validated using an independent CAPS-based assay. Finally, automated qPCR screening of a wheat HII population to identify mutants for each TaPFT1 homoeologue was performed and identified mutants validated using the CAPS-based assay. After developing and validating this method using TaPFT1, we applied it to the detection of homoeologue deletions of three additional target genes (TaNFXL1, TaCesA8 and TaPLD1) and successfully identified multiple deletions for each within the HII population (Table 2).

We used patented Applied Biosystems MGB-NFQ TaqMan SNP-detection probes for all assays (see Materials and Methods). We chose this supplier because ABI provided a flexible custom probe design service and assistance with the design of these assays to our specific criteria. However, a range of other SNP-detection
fluorogenic qPCR probes are available from other suppliers, and these may also be adaptable to this assay.

As outlined above, using the assay developed we were able to determine the chromosomal location of TaPFT1-A by screening nullisomic-tetrasomic lines of the wheat genotype Chinese Spring. We subsequently screened Chinese Spring deletion lines to narrow down its genomic location to a region between FL 0.11 and FL 0.35 on chromosome 5A, close to the centromere region (data not shown). Similarly, using nullisomic-tetrasomic lines of Chinese Spring, TaPFT1-B and TaPFT1-D were shown to reside on chromosomes 5B and 5D, respectively. Lines with 5B and 5D chromosome deletions that are large enough to exclude these loci based on the location of TaPFT1-A were not available, and all screened 5B and 5D deletion lines possessed the respective TaPFT1 homoeologues. However, from the results of the nullisomic-tetrasomic lines in conjunction with the largest 5B and 5D long and short arm deletions screened, we could putatively assign TaPFT1-B to a location between 5BS FL 0.13 and 5BL FL 0.52; and TaPFT1-D to between 5DS FL 0.22 and 5DL FL 0.60, and these locations are consistent with the location determined for TaPFT1-A. However, since none of the smaller 5B and 5D deletion lines screened lacked the respective copies, these finer chromosomal locations of TaPFT1-B and TaPFT1-D will require further definition in the future.

The location of TaPFT1 homoeologues on their respective homoeologous chromosomes was confirmed using a TaPFT1 CAPS assay (Figure 5B) designed for validation of the probe-based assay. The chromosomal location of TaPFT1 in wheat is consistent with the demonstrated location of OsPFT1 on chromosome 9 (LOC_Os09g13610 within the 'Rice Genome Annotation Project') which is syntenic to wheat chromosome 5 [42]. Additionally, blasting TaPFT1 against the Phytozome v5.0 database http://www.phytozome.net has revealed that putative Sorghum bicolor SbPFT1 and Brachypodium distachyon BsPFT1 are located on chromosomes 2 and 4, respectively - Sorghum chromosome 2 and Brachypodium chromosome 4 are indeed syntenic to wheat chromosome 5 [42,43]. However, our results assigning TaPFT1 on wheat chromosome 5 conflict with a location of a 554 bp putative TaPFT1 EST (accession no. BF472999), which shows approximately 97% homology to TaPFT1 sequence that we obtained, on chromosome 7DS in the GrainGenes 2.0 database http://wheat.pw.usda.gov/GG2/index.shtml. One possible explanation for this discrepancy might be the existence of a duplicate TaPFT1 locus. However, the results of our sequencing of TaPFT1 clearly showed the presence of only three distinct sequences in the hexaploid wheat genotype we used, arguing against the presence of such a duplicate locus. Another possibility could be that a translocation of a TaPFT1 locus has occurred in the specific genotype used in mapping. Another, more probable, explanation is that this EST has been incorrectly mapped in the GrainGenes database.

Prior to the development of this technique, screening of the HII lines used in this study for homoeologue deletions of a number of candidate genes had been performed using CAPS assays similar to that performed in this study for validation purposes. The key disadvantages of CAPS screening include: difficulty in identifying sufficient polymorphism for design of the assay, and high labour and time costs of performing large-scale screening. By exploiting probes designed to effectively discriminate between sequences differing by as little as a single base pair, this new assay requires only the most minimal polymorphism between homoeologues. The minimum requirement for successful design of the specific probes are the presence of individual SNPs specific for each homoeologue occurring within a region of suitable length for a qPCR amplicon (e.g. <400 base pairs), that can be amplified from each homoeologue in a single reaction. Where possible, a single primer pair has
been used for an individual assay. However, polymorphism in an otherwise desirable site for primer design can be accommodated via the multiplexing of primers showing identity to each of the polymorphic sequences, as for the TaPLD assay (Table 2). With our previous screening using CAPS, a single researcher could process approximately 200 samples per day. In contrast, this new assay allows us to screen up to 384 individual samples (192 individual samples in duplicate) in approximately 2 hours. The time taken to perform the qPCR assay is the limiting factor for screening - a robotic set up of individual plates takes approximately one hour. Using our equipment that allows pre-prepared optical plates to be stacked and continuously loaded and analysed, this theoretically allows for the processing of approximately twelve 384-well plates, corresponding to up to approximately 2000 duplicated samples, in a 24 hour period, in an automated manner, by a single researcher.

It should be noted that, as for genome-specific PCR and CAPS-based deletion screening techniques, this assay is not suitable for the detection of hemizygous individuals (those in which a specific locus is missing from one homologous chromosome but intact within...
the other) and is not amenable to sample pooling due to its reliance on detection of the lack of specific target sequence. Although the ability to detect individuals hemizygous for a target locus would be advantageous, the accurate identification of hemizygous lines is inherently challenging as the target sequence remains present in these lines, albeit at half the concentration. Whilst it is theoretically possible to discriminate copy number of a target locus using qPCR, we have found that when dealing with variation in DNA sample concentration and quality that can be present in any large collection of samples such as those screened here, hemizygous individuals may not be identified with confidence using this method.

We chose to screen samples in duplicate to provide additional validation of putative mutants, and to cater for failure of a single reaction due to technical error. During the process of screening for homoeologous deletions of the four target genes, we have assessed approximate frequencies of conflicting screening results between duplicate reactions. Technical errors leading to one successful and one failed reaction are indeed very low (< 1 per 96 qPCR reactions); whilst mutant/non-mutant fluorescence profiles, indicative of false positive or negative results, are absent where CT values are < 30 and very rare for DNA samples with low concentrations yielding CT values >35 (< 1 per 384 qPCR reactions). Therefore, duplication for deletion screening using this

| gene name | sequence length (bp) | GenBank accessions | primer sequences1 ; amplicon size1 | probe sequences2 assay conditions3 | samples screened; mutations detected |
|-----------|----------------------|--------------------|-----------------------------------|-----------------------------------|-------------------------------------|
| TaNXL1    | ~630                 | HQS95068 - HQS95070| FNXL1gsprb GGCACACCACTCCATCATG RNXL1gsprb GGCTGGTTGCATCTGATATC | NFXL1prb1FAM AACACATTCTCTTGACTG NFXL1prb2VIC AACGCACTTCGTGACTG NFXL1prb3NED ACATTCTCTGATACACA | FR primers: 1000 nM each Probes: 50 nM FAM, 100 nM VIC, 100 nM NED 20 μL reaction: 18 μL stock; 2 μL DNA sample 3-step reaction: 95°C 15 s denaturation, 55°C 30 s annealing, 60°C 30 s extension - 45 cycles |
| TaCesA8   | ~370                 | HQS95065 - HQS95067| FCesA8gsprb CCCATGGCGAAGACACCT RCesA8gsprb AGTGAACCGACGTGCTAC | CesA8prb1FAM GTCATAACCCGCTATTGA CesA8prb2VIC CGGACCAGGTGAG CesA8prb3NED ACGAGGTGAGAACC | FR primers: 1000 nM each Probes: 50 nM FAM, 100 nM VIC, 100 nM NED 20 μL reaction: 18 μL stock; 2 μL DNA sample 3-step reaction: 95°C 15 s denaturation, 55°C 30 s annealing, 60°C 30 s extension - 45 cycles |
| TaPLDβ1   | ~620                 | HQS95071 - HQS95073| FPLDB1gsprb TTTGATCTGTCCTTGATGTTATG R1PLDB1gsprb GCCGTAGTGGAGCGAGTC TC R2.3PLDB1gsprb GGTGTAGTGGAGCGAGTC | PLDB1prb1FAM TTGCTTAGAAGCTGC GG PLDB1prb2NED CTAGAACCAGCTGTTT PLDB1prb3VIC TAGAACCAGCTGTTT | FR primers: 1000 nM F1 primer, 666 nM R2.3 primer, 333 nM R1 primer Probes: 50 nM FAM, 100 nM VIC, 150 nM NED 11 μL reaction: 9 μL stock; 2 μL DNA sample 3-step reaction: 95°C 15 s denaturation, 55°C 30 s annealing, 60°C 30 s extension - 45 cycles |

1Approximate; some variation between homoeologues was present 2All primer and probe sequences are given in 5'-3' directions 3All assays run in 'Standard Mode' on an ABI 7900HT (see Materials and Methods); all thermal cycling commences with 50°C 2 mins, 95°C 10 mins
method is arguably unnecessary, especially since downstream validation of indexed seed will be required for any identified mutant lines. By eliminating duplication, throughput of this method could be doubled, and cost per sample halved.

Another technology that could be potentially explored for deletion screening in polyploid plant species is automated single-strand conformation polymorphism analysis (SSCP)-based capillary electrophoresis. Theoretically, this technology could allow for the detection of homoeologue-specific target gene deletions. However, the logistics of such an application for this method are unknown, and a recent protocol for mutation screening using this technique indicates that throughput for sample screening (a maximum of 768 samples over three days) [44] is unlikely to be capable of matching that of the presented method (approximately 6000 duplicated samples in the same time period).

Recently, a modified TILLING protocol for the screening of hexaploid wheat for mutations has been developed [15]. This method appears to be an extremely powerful tool for screening to detect point mutations and small deletions simultaneously within homoeologues of mutagenized wheat lines, such as those generated using EMS [6]. The technique outlined here is not appropriate for the screening of homoeologues for non-specific point mutations and small deletions. Rather, it is useful for detection of presence or absence of homoeologous copies of candidate genes in a mutagenized population within which lines possess frequent whole-gene deletions such as the HII population used in this study. For this purpose, this method has significant advantages over any other currently available technique. In addition, HII lines are considered non-GM; therefore, mutants with beneficial characteristics can be integrated into crop improvement programs. However, as for lines generated using chemical mutagenesis, HII lines can possess ‘background’ mutations. For reverse genetics approaches to crop improvement using this method in conjunction with a HII population, identified lines with beneficial characteristics can be relatively easily ‘purified’ by successive rounds of backcrossing and re-screening to remove the majority of such background mutations.

As reported here, our design and optimization of this technique has been performed in hexaploid wheat, which globally is the most significant polyploid crop plant. Like wheat, many important plant species including cotton, canola and potato (tetraploid), triticale and oat (hexaploid), strawberry (octoploid), and sugarcane (5 n-14 n [45]) are polyploids. Therefore, this technique has the potential to be adapted to many other polyploid species. In addition to screening for homoeologues in other polyploid species, this assay would be readily adaptable to the screening for duplicated gene copies or closely-related gene family members within diploid species so long as SNPs that allow the design of gene-specific fluorescent probes exist in the sequences to be identified. Whole-genome sequencing efforts are currently underway in many polyploid and diploid crop plants including wheat. The availability of assembled whole-genome sequence information would facilitate the efficient design of assays for virtually any gene/s using this method.

Conclusions
Here we present a method for automated, high-throughput screening to detect and distinguish individual homoeologues of any wheat gene. We designed this method for the purpose of identifying homoeologous deletions of target genes within a resource of HII mutagenized wheat lines featuring frequent whole-gene deletions. This method is potentially applicable to the identification of homoeologous copies of a target gene within any polyploid species. It may also have utility for the identification of highly homologous genes (e.g. duplicated genes or gene family members) within diploid species.

Methods
Plant material
The commercial hexaploid bread wheat cultivar Chara (AWB Seeds Ltd., Australia) was used for characterisation of target gene sequences and for the isolation of homoeologous deletion mutants in these genes. Heavy ion irradiation of grain of cv. Chara was conducted by the RIKEN RI-beam facility in Japan [46] using a Neon ion beam at 50 Gy. Irradiated grain was germinated and selfed to generate an M2 population comprising approximately 20000 lines and leaf material sampled for DNA extraction. Nullisomic-tetrasomic wheat lines [31,32] and deletion wheat lines [33] derived from the cultivar Chinese Spring were used to determine the chromosomal location of the TaPFT1 gene. These were kindly provided by Drs. Evans Lagudah and Chunji Liu of CSIRO Plant Industry and are maintained at the USDA-Agricultural Research Service, University of Missouri, Columbia, MO, USA. All plants were grown under glasshouse conditions in either Canberra or Brisbane, Australia.

DNA Extraction
DNA was extracted from leaf tissue using a Qiagen DNeasy Plant Mini Kit (Qiagen GmbH) in conjunction with a QIAcube (Qiagen GmbH) apparatus for all samples.

Design of primers and probes
The Applied Biosystems (Applied Biosystems, Foster City, CA) custom probe design service was used to assist
in the design of primers and patented minor-groove-binding, non-fluorescent quencher (MGB-NFQ) TaqMan probes. Homoeologous TaPFT1 sequences used in the design of the assay with highlighted primer and probe sequences are shown in Figure 2A. The position of the region within TaPFT1 is highlighted in Figure 1. Primer and probe sequences for the TaNFXL, TaCesA8, and TaPLDB1 assays are given in Table 2.

Sequencing of target genes
Primers for amplification of genomic regions of TaPFT1, TaNFXL1, TaCesA8 and TaPLDB1 were designed using Primer Premier 5.1 (Premier Biosoft International, Palo Alto, CA). PCR products were amplified from genomic Chara DNA using Phusion High-Fidelity DNA polymerase (Finnzymes Oy, Keilaranta 16 A, 02150 Espoo, Finland) with a manufacturer-reported error rate of 4.4 x 10^-7. PCR products were cloned using a TOPO TA cloning kit (Invitrogen, Carlsbad, CA, USA). Plasmid preparations were prepared using a QIAprep Spin Miniprep kit (Qiagen GmbH) in conjunction with a QIAcube (Qiagen GmbH) apparatus. The Australian Genome Research Facility (AGRF) plasmid sequencing service (AGRF, Brisbane, QLD, Australia) was used to obtain cloned fragment sequence using forward and reverse M13 primers. Sequence was analysed using Sequencher Version 4.9 Software (Gene Codes Corporation, Ann Arbor, MI, USA). For TaPFT1, sequences labelled TaPFT1a(1-3) - TaPFT1e(1-3) (see Results) were deposited in GenBank (accession numbers HM561279 - HM561293). Labelling of sequences 1-3 was arbitrary and not indicative of origin in terms of progenitor genome. However, genomic origin of the TaPFT1b and TaPFT1c sequences used in the design of the probe and CAPS assays respectively have subsequently been determined, as outlined in Figure 1 and 2, by screening of nullisomic-tetrasomic ‘Chinese Spring’ lines. For TaNFXL1, TaCesA8, and TaPLDB1, sequences labelled 1-3 prefixed with gene name were deposited in GenBank. Again, numerical labelling of sequences was arbitrary for each gene. Accession numbers are given in Table 2.

Optimization of assay for TaPFT1
Primers were tested for PCR specificity using Chara DNA template in conjunction with TaqMan Universal PCR Mastermix (Roche Diagnostics GmbH, Mannheim, Germany). PCR was performed using an Applied Biosystems 9600 thermal cycler apparatus (Applied Biosystems, Foster City, CA). PCR conditions were an initial denaturation step at 95°C for 10 minutes, then 45 cycles of a 15 sec, 95°C denaturation step followed by a 60 sec, 60°C annealing/extension step. 5 μL PCR product was visualized on a 1.4% agarose gel stained with Gel Red (Biotum Inc., Hayward, CA) against 1 μL of Invitrogen 1 kb+ DNA ladder (Invitrogen, Carlsbad, CA, USA). A single band of the desired size was observed.

The multiplexed qPCR assay was optimized using Chara DNA template. Final concentrations of forward and reverse primers remained standard at 1000 nM each. All combinations of final concentrations of 50, 100, and 150 nM for the three probes were assessed. Primer/probe mixes were combined with TaqMan Universal PCR mix (final concentration 1×) and water to produce stocks. 9 μL of stock was added to 1 μL 100 ng/μL Chara DNA, or water as a negative control. Four replicate optimization reactions in conjunction with a negative control were set up for each probe concentration combination. All reactions were set up in a single MicroAmp 384-well optical plate (Applied Biosystems, Foster City, CA) and sealed with a MicroAmp optical adhesive cover (Applied Biosystems). qPCR was performed for these samples using an Applied Biosystems 7900HT qPCR apparatus (Applied Biosystems, Foster City, CA). Thermal cycler was run in ‘9600 Emulation’ mode. Fluorescence detection was in ‘Absolute Quantification’ mode using three detection channels: Fam-NonFluorescent; Ned-NonFluorescent; and Vic-NonFluorescent. Individual detection channels corresponded to fluorescence produced by individual probes. Optimum fluorescence for all probes in the multiplexed assay was obtained using final concentrations of 50 nM for the ‘FAM’ and ‘VIC’ labelled probes and 100 nM for the ‘NED’ labelled probe. Under these reaction conditions efficient increase of fluorescence from all probes was observed, with C_T values of approximately 25 cycles for all probes (Figure 3).

In order to assess the specificity of probe fluorescence, 1/10000 dilutions of plasmid preparations, determined by sequencing to harbour cloned PCR products corresponding to regions of homoeologues 1′, 2′, and 3′ targeted by the assay, were prepared. Four replicate reactions for each of five templates: diluted clone A (homoeologue 1), diluted clone B (homoeologue 2), diluted clone C (homoeologue 3), Chara (positive control), and water (negative control) were prepared in a 384-well optical plate. qPCR was performed as above. For all replicates using specific diluted clone templates, efficient fluorescence increase was observed from the corresponding specific probe (i.e. diluted clone 1 template: efficient ‘FAM’ fluorescence; diluted clone 2 template: efficient ‘NED’ fluorescence; diluted clone 3 template: efficient ‘VIC’ fluorescence), but not from the non-specific probes. This indicated that the probe fluorescence was highly specific; non-specific fluorescence was low, or absent, even where non-specific homoeologous sequence was in high concentration and not in competition with the specific homoeologous sequence.
Optimization of assays for other target genes followed the same basic steps as outlined above i.e. 1. assess primer specificity; 2. optimize multiplexed probe assay against Chara DNA template; and 3. assess specificity of probe fluorescence using diluted cloned homoeologue sequence. For the TaNFXL1 and TaCesA8 assays, reaction volumes were increased to 20 µL (18 µL stock; 2 µL DNA sample) for optimum results. Details of optimized assays are given in Table 2.

Assessment of limit of detection
To assess the limit of detection of the method in terms of DNA concentration in a sample, serial 1:5 dilutions of a Chara DNA sample were prepared such that resulting samples had concentrations of 91 ng/µL (undiluted); 18.2 ng/µL; 3.7 ng/µL; 0.7 ng/µL; 0.14 ng/µL. 2 µL volumes of these samples were then assayed using the TaPFT1 probe-based method, and the TaPFT1 CAPS method. Both methods showed consistent results (i.e. the same profiles in three independent reactions) with as little as 0.7 ng/µL DNA concentration.

High-throughput, automated screening of HII mutant lines to identify individuals lacking homoeologous copies of target genes
The optimized probe method was used for the screening of 2500-4500 individual HII-generated Chara mutant lines per gene in order to identify mutants lacking homoeologues of target genes. DNA from individual M2 mutant lines arranged in indexed 96-well plate format of primers and probes, and water was combined with Universal Master Mix (Roche), optimized concentrations of concentrations in a sample, serial 1:5 dilutions of a Chara DNA sample were prepared such that resulting samples had concentrations of 91 ng/µL (undiluted); 20.2 ng/µL; 3.7 ng/µL; 0.7 ng/µL; 0.14 ng/µL. 2 µL volumes of these samples were then assayed using the TaPFT1 probe-based method, and the TaPFT1 CAPS method. Both methods showed consistent results (i.e. the same profiles in three independent reactions) with as little as 0.7 ng/µL DNA concentration.

'TCAPS' screen for validation of mutants detected by TaPFT1 probe screen
A region of TaPFT1 with relatively high polymorphism between homoeologues, suitable for the design of a CAPS screen [47], was identified from sequence obtained for TaPFT1 (Figure 2B). Primers to amplify this region from all homoeologues non-specifically were designed using Primer Premier 5.1 (Premier Biosoft International, Palo Alto, CA). A single restriction enzyme producing clearly distinguishable banding patterns upon cleavage of each homoeologue was identified using the New England Biolabs 'NEBcutter V2.0' application [48] (Figure 5A). The CAPS screen was optimized using Chara DNA. Upon digestion of the amplified product, the expected banding pattern (as predicted by NEBcutter V2.0) was observed. The optimized CAPS screen was performed on Chinese Spring DNA, in conjunction with nullisomic-tetrasomic DNA determined by the probe-screen to lack homoeologous copies of TaPFT1. The banding patterns for Chinese Spring, and Null5A, Null 5B and Null 5D (shown to lack the respective homoeologous copies of TaPFT1) samples were as predicted by 'Virtual Digest' (Figure 5B). DNA from seedlings corresponding to each specific mutant line identified as lacking a PFT1 homoeologue was assessed using the optimized CAPS screen described above.

Southern blotting of TaPFT1 CAPS fragments
DNA fragments were transferred to an Amersham Hybond-XL (GE Healthcare Life Sciences) nylon membrane using the 'Neutral transfer protocol' as outlined in the Amersham Hybond-XL product booklet. Fragments were hybridised with a labelled TaPFT1 fragment amplified from cloned TaPFT1 fragment ‘B’ (Figure 1) plasmid preparation using TaPFT1 specific forward and reverse primers and radiolabelled with [alpha-<sup>32</sup>P] dCTP using an Amersham Megaprime DNA labelling kit (GE Healthcare Life Sciences). Hybridisation was performed overnight at 65°C and the blot washed in high stringency according to instructions given by the manufacturer. The blot was then exposed to an X-ray film and developed using standard procedures.

Additional material

Additional file 1: Figure S1. Diagrammatic overview of all partial sequences (TaPFT1a - TaPFT1e) obtained for wheat TaPFT1 homoeologues.

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Author details
1CSIRO Plant Industry, 306 Carmody Road, St Lucia, QLD 4067, Australia. 2CSIRO Plant Industry, GPO Box 1600, Canberra ACT 2601, Australia. 3CSIRO Food Futures National Research Flagship, PO Box 93, North Ryde 1670, NSW, Australia.

Authors’ contributions
ZL and MMH developed the HII wheat mutant population resource and provided DNA samples. TLF designed and optimized the method, performed
mutation screening, and assisted in writing the paper. KK and JMM assisted in design and coordination of the project and in writing the paper. All authors read and approved the final manuscript.

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