1	Identification and comparison of individual chromosomes of three Hordeum
2	chilense accessions, Hordeum vulgare and Triticum aestivum by FISH
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24 Abstract

25 Karyotypes of three accessions of Hordeum chilense (H1, H16 and H7), Hordeum vulgare and Triticum 26 aestivum were characterized by physical mapping of several repetitive sequences. A total of fourteen 27 repetitive sequences were used as probes for fluorescence in situ hybridization (FISH) with the aim of 28 identifying inter- and intra-species polymorphisms. The (AG)12 and 4P6 probes only produced 29 hybridization signals in wheat, the BAC7 probe only hybridized to the centromeric region of *H. vulgare*, 30 and the pSc119.2 probe hybridized to both wheat and H. chilense, but not to H. vulgare. The remaining 31 repetitive sequences used in this study produced a hybridization signal in all the genotypes. Probes pAs1, 32 pTa535, pTa71, CCS1 and CRW were much conserved, showing no significant polymorphism among the 33 genotypes studied. Probes GAA, (AAC)₅, (CTA)₅, HvT01 and pTa794 produced the most different 34 hybridization pattern. We identified large polymorphisms in the three accessions of *H. chilense* studied, 35 supporting the proposal of the existence of different groups inside *H. chilense* species. The set of probes 36 described in this work allowed the identification of every single chromosome in all three species, providing 37 a complete cytogenetic karyotype of H. chilense, H. vulgare and T. aestivum chromosomes, useful in wheat 38 and tritordeum breeding programs.

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40 Key words: Hordeum, Triticum aestivum, tritordeum, FISH, repetitive sequences

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50 Introduction

51 Hordeum chilense Roem. et Schultz. $(2n = 2x = 14; H^{ch} H^{ch})$ is a diploid wild barley, native of Chile and Argentina (von Bothmer et al. 1995), which shows very useful traits for wheat breeding, including drought 52 53 and salt tolerance (Gallardo and Fereres 1980; Forster et al. 1990), resistance to pests and diseases (Martín 54 et al. 1996) and high seed carotenoid content (Atienza et al. 2004). Several substitution and addition lines 55 have been developed to transfer these traits into wheat (Miller et al. 1982; Fernández and Jouve 1988), and 56 recently, some accessions of *H. chilense* have been used to produce male sterility in wheat with the aim of 57 establishing a wheat hybrid system (Martín et al. 2008; Martín et al. 2010; Castillo et al. 2014). In addition, 58 a major interest in this species relates to its high crossability with other members of the Triticeae tribe (von 59 Bothmer et al. 1986: Martín et al. 1998). Evidence of this is the obtaining of fertile amphiploids named 60 tritordeum (XTritordeum Ascherson et Graebner), which were obtained after chromosome doubling of 61 hybrids between H. chilense and tetraploid and hexaploid wheats (Martín and Sánchez-Monge Laguna, 62 1982; Padilla and Martín 1983; Martín et al. 1999). In addition to H. chilense, other Hordeum species have 63 been hybridized with wheat; however, the only amphiploid showing enough fertility to be considered as a possible crop is ×Tritordeum martinii (HchHchAABB) (Pujadas Salvá 2016), which resulted from 64 65 interspecific crosses between H. chilense and cultivated durum wheat. Today, Tritordeum martinii is grown 66 in Spain, France, and Italy, being the only synthetic amphiploid crop commercialized for human 67 consumption.

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Hordeum chilense is an extremely variable species, which can be found in many contrasting environments (Tobes et al. 1995; Giménez et al. 1997). It shows a very wide range of variation of morphological characters (von Bothmer et al. 1980, 1995), different levels of avoidance to barley leaf rust (Rubiales and Niks 1992, 1996) and it even differs at the cytoplasmic level (Atienza et al. 2007; Martín et al. 2009). In 2001, a study concluded that *H. chilense* consists of at least three morphologically and genetically distinct subspecific taxa (Vaz Patto et al. 2001). However, not much work has been done to identify or define further these three subspecific taxa or groups.

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Hordeum chilense, and the genus Hordeum in general, has been characterized by FISH using several
 repetitive sequences. However, most of the repetitive probes used in this genus were aimed at identifying
 the 7 pairs of Hordeum chromosomes, and more repetitive sequences are required to identify individual

80 chromosome arms or smaller regions introgressed into wheat, which ultimately, is the goal in most81 introgression breeding programs.

In this work, we use fourteen repetitive sequences, for the identification and comparison by FISH, of all individual chromosomes of *H. chilense*, *H. vulgare* and *T. aestivum*. Furthermore, we increase the knowledge of the karyotype of the different *H. chilense* groups described, by using these probes in the analysis of three *H. chilense* accessions (H1, H16 and H7), each one of them belonging to one of the three different groups.

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88 Materials and methods

89 Plant material

Hordeum vulgare cv. Betzes, *Tritordeum martinii* "HT27" and three accessions of *Hordeum chilense*(H1, H16 and H7) were kindly provided by Prof. Antonio Martín from the Instituto de Agricultura
Sostenible (CSIC, Spain). *Hordeum chilense* accessions H1, H16 and H7 belong to the group I, II and III
respectively, described by Vaz Patto et al. (2001). *Triticum aestivum* cv. Chinese Spring was also used in
this work.

95 Fluorescent in situ hybridization (FISH)

96 Chromosome preparation for FISH

97 Seeds of each genotype used in this study were germinated on wet filter paper in the dark for 5 days at 98 4°C, followed by a period of 24h-48h at 25°C under constant light. Root tips were obtained from these 99 seedlings and occasionally, from adult plants grown in pots in a controlled environmental room (16/8 h, 100 light/dark photoperiod at 20 °C day and 15°C night, with 70% humidity). Preparation of chromosome 101 spreads was done following Kato et al. (2004) and King et al. (2017) with slight modifications. These 102 modifications were: roots were mainly excised from seedlings, although occasionally, adult plants were 103 also used in the obtaining of chromosomes spreads. Also, the percentage of cellulase was increased from 104 2% to 4%, being the final enzymatic cocktail: 1% pectolyase Y23 and 4% cellulose Onozuka R-10 (Yakult 105 Pharmaceutical, Tokyo) solution in 1x citrate buffer.

106 Development of probes and labelling

107 Fourteen repetitive sequences were used in this work (Table 1). All of them were amplified by PCR and 108 the primer sequences and PCR conditions are described in Table 1. Three different polymerase enzymes 109 were used: BIOTOOLS[™] DNA polymerase (Biotools S.A., Madrid, Spain); KAPA3G plant polymerase 110 (VWR International GmbH, Erlangen, Germany); and MyFi DNA polymerase (Bioline USA, Taunton, 111 MA) (Table 1). PCR was performed in 50 µl reaction mixtures according to the manufacturer's instructions 112 of each polymerase, and PCR products were resolved on 2% agarose gels in 1xTBE, stained with ethidium 113 bromide and visualized under UV light. The PCR products were purified using a Qiagen kit (Qiagen, 114 Hilden, Germany) following the manufacturer's instructions. The purified PCR products were labelled with 115 either biotin-16-dUTP or digoxigenin-11-dUTP, using the Biotin-nick translation mix or the DIG-nick 116 translation mix, respectively (Sigma, St. Louis, MO, USA) according to the manufacturer's instructions. 117 Nick translation was performed in a PCR machine at 15°C for 90 min followed by a final step at 68°C for 118 10 min. Hordeum chilense was also used to re-probe the samples after FISH with the repetitive sequences 119 in order to identify the H. chilense chromosomes in the tritordeum line (HT27). Hordeum chilense DNA 120 was extracted from leaves of young plants following the CTAB method described in Murray and Thomson 121 (1980).

122 FISH and GISH experiments

123 The FISH protocol used in this work was a combination of two protocols previously described in 124 Cabrera et al. (2002) and King et al. (2017). The FISH procedure was performed over two days. Unless 125 indicated otherwise, washes were carried out at room temperature. First day: Slides were crosslinked in an 126 UV Cross linker at 0.125 Joules for 30 secs (twice). The hybridization mixture consisted of 50% formamide, 127 10% dextran sulfate, 2x saline sodium citrate (SSC), 0.125% sodium dodecyl sulfate (SDS) and 0.1 mg of 128 salmon sperm DNA. The concentration of each probe in the hybridization mixture is described in Table 1. 129 The hybridization mixture was denatured for 8 min at 80 °C and cooled on ice for 5 min. A 40 µl-aliquot 130 of the hybridization mixture was added to the cross-linked samples and a cover-slip applied. Slides were 131 incubated in an in-situ PCR thermal cycler (Leica Microsystems[™] ThermoBrite[™], Leica Biosystems, 132 Wetzlar, Germany) at 80°C for 7 min to denature the chromosomes. The hybridization was carried out overnight (20-24h) at 37 °C in a humid chamber. Second day: slides were washed in 2xSSC twice for 5 133 134 min at 37°C, in 1xSSC twice for 5 min and in TNT (0.1 mol/L Trix-HCL, 0.15 mol/L NaCL, 0.05% Tween-20) for 5 min. Slides were then blocked in 50% (w/v) dried skimmed milk in TNT for 20 min at 37°C and 135 136 washed in TNT for 2 min. For the detection of biotin and digoxigenin labelled probes, slides were incubated at 37°C for 45 min with streptavidin-CY3 (Sigma) and antidigoxigenin-FITC (Sigma) in 1x phosphate
buffered saline (PBS), respectively (Table 1). Slides were then washed in TNT for 5 min and dehydrated
in 70% and 100% ethanol for 1 min. After counter-staining with 4', 6-diamidino-2-phenylindol (DAPI) for
5 min, slides were washed in water for 5 min, dehydrated again and mounted in Vectashield (Vector
Laboratories, Burlingame, CA, USA). In tritordeum "HT27", all chromosome spreads were re-hybridized
following the reprobing method of Heslop-Harrison et al. (1992), in order to identify the *H. chilense*chromosomes in the wheat background.

144 Image acquisition and chromosome identification

145 Hybridization signals were examined using a Leica DM5500B microscope equipped with a Hamamatsu

146 ORCA-FLASH4.0 camera and controlled by Leica LAS X software v2.0. Digital images were processed

using Adobe Photoshop CS5 (Adobe Systems Incorporated, USA) extended version 12.0×64 .

The ideogram for H1, H16, H7, *H. vulgare* and *T. aestivum* chromosomes was based on the
hybridization patterns of the probes used in this work and the morphology of chromosomes previously
described (Cabrera et al. 1995; Pedersen and Langridge 1997; Prieto et al. 2004; Kato 2011; Szakács et al.
2013; Komuro et al. 2013; Tang et al. 2014).

152

153 Results

154 Repetitive sequences hybridized to mostly terminal and interstitial regions: pAs1 and pTa-535

The pAs1 probe is a repetitive DNA sequence isolated from *Aegilops tauschii* Coss. (formerly known
as *Ae. squarrosa* L.) (Rayburn and Gill 1986). The pTa-535 is a 342-bp tandem repeat isolated from *T*. *aestivum* L. (Komuro et al. 2013).

Both probes show similar hybridization patterns in all the *Hordeum* analysed in this work, H1, H16, H7 and *H. vulgare*, with all the chromosomes showing signals in both arms (Fig. 1). The three accessions of *H. chilense* H1, H16 and H7 show no major differences in their hybridization patterns (Fig. 1). The lowest and weakest number of hybridization signals per chromosome was observed in *H. vulgare* (Fig. 1). Both pAs1 and pTa-535 predominantly hybridized to the telomeric and subtelomeric regions, with some interstitial and centromeric signals in some chromosomes (Fig. 1). In H1, H16 and H7, chromosomes 1H^{ch} and 4H^{ch} exhibited weaker signals than the rest of the chromosomes; while in *H. vulgare*, chromosomes

 $165 \qquad 3H^{v},\,4H^{v} \text{ and } 5H^{v} \text{ exhibited the weakest signals (Fig. 1)}.$

In wheat, both probes mainly hybridized to both chromosome arms in all D genome chromosomes as previously described (Rayburn and Gill 1986; Komuro et al. 2013). Signals were also predominantly telomeric and subtelomeric, as was observed in the *Hordeum* analysed. Some chromosomes from the A and B-genomes occasionally hybridized to both probes, but these signals were weak and unsteady, so only Dgenome chromosomes were identified in this work.

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172 Repetitive sequences hybridized to terminal regions: pSc119.2 and HvT01

The pSc119.2 probe is a repetitive sequence containing 120 bp from *Secale cereal* L. (Bedbrook et al.
174 1980). The HvT01 is a subtelomeric sequence from *H. vulgare* L. (Belostotsky and Ananiev 1990).
175 Hybridization signals with both pSc119.2 and HvT01 were polymorphic in all plant material analyzed in
176 this study (Figs. 2, 5).

177 In *H. chilense* H1, six pairs and seven pairs of chromosomes were hybridized to pSc119.2 and HvT01, respectively (Fig. 2). Probe pSc119.2 did not hybridize on chromosome 3H^{ch}, and the signals produced on 178 179 chromosome 7H^{ch} were weak (Fig. 2). The pSc119.2 signals were detected on the short arm of 1H^{ch}, 2H^{ch}, 180 5H^{ch} and 7H^{ch} (1H^{ch}S, 2H^{ch}S, 5H^{ch}S and 7H^{ch}S), on the long arm of 6H^{ch} (6H^{ch}L) and on both arms of 4H^{ch} 181 (Fig. 2). Probe HvT01 was detected on 2H^{ch}S, 4H^{ch}S, 5H^{ch}S, 6H^{ch}L and 7H^{ch}S, and on both arms of 1H^{ch} 182 and 3H^{ch} (Fig. 2). In *H. chilense* H16, four pairs of chromosomes were labelled with pSc119.2 and HvT01 183 (Fig. 2). The pSc119.2 signals were detected on 1H^{ch}S, 5H^{ch}S, 6H^{ch}L and on both arms of 4H^{ch} (Fig. 2). 184 The HvT01 signals were detected on 3H^{ch}S, 4H^{ch}S, 6H^{ch}L and on both arms of 5H^{ch} (Fig. 2). In H. chilense 185 H7, both repetitive sequences (pSc119.2 and HvT01) were detected on 3 pairs of chromosomes (Fig. 2). 186 The pSc119.2 signals were observed on 5H^{ch}S and on both arms of 1H^{ch} and 4H^{ch} (Fig. 2). The HvT01 187 signals were detected on 3H^{ch}S, 4H^{ch}S and 5H^{ch}S (Fig. 2). The HvT01 signals obtained in H1 and H7 agreed 188 with the results of Prieto et al. (2004).

In *H. vulgare*, no hybridization signal was detected using the pSc119.2 probe (Fig. 2), which agreed
with previous studies (Gupta et al. 1989). However, all pairs of chromosomes were hybridized to HvT01
on both chromosome arms, except 2H^vS (Fig. 2).

192	In wheat, thirteen pairs of chromosomes were hybridized to the pSc119.2 (Fig. 5). Signals were detected
193	on all B-genome chromosomes and on chromosomes 4AL, 5AS, 2DS, 3DS, 4DS and 5DS (Fig. 5). All the
194	pSc119.2 signals were detected at the telomeric and subtelomeric regions (Fig. 5) as observed in the genus
195	Hordeum. Probe HvT01 did not produce any signal in wheat. Occasionally, two pairs of chromosomes
196	showed a very weak signal, but since the results were not consistent, they were not considered in this work
197	(Fig. 5).

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199 Repetitive sequences hybridized mostly to interstitial, centromeric and pericentromeric regions: 200 GAA, (AAC)5, (CTA)5, 4P6 and (AG)12

The GAA probe is a GAA-satellite sequence from wheat and barley (Dennis et al. 1980; Pedersen et al.
1996). The (AAC)₅ probe is a tri-nucleotide repeat from *T. aestivum* L. (Cuadrado et al. 2008). The (CTA)₅
probe is a tri-nucleotide repeat from *T. aestivum* L., which has been used for the first time in this work. The
4P6 probe is a tandem repeat BAC clone from *Ae. tauschii* Coss. (Zhang et al. 2004). The (AG)₁₂ probe is
a di-nucleotide repeat from *T. aestivum* L. (Cuadrado et al. 2008).

206 The GAA sequence was abundant in all the analysed species, allowing for the identification of all 207 Hordeum chromosomes and all the wheat B-genome chromosomes (Figs. 3,5). This probe predominantly 208 hybridized to interstitial and pericentromeric regions, with some distal signals in some chromosomes. In H. 209 chilense H1, several pericentromeric signals were detected on chromosomes 3H^{ch} and 7H^{ch}, and several 210 interstitial signals on 4H^{ch}S and 2H^{ch}S. Chromosome 1H^{ch}L was the only chromosome showing a strong 211 terminal signal. Chromosome 5H^{ch}S and 7H^{ch}S showed occasionally a weak interstitial and telomeric signal, 212 respectively. In *H. chilense* H16, the hybridization pattern was slightly different to H1. Chromosomes 6H^{ch} 213 and 7H^{ch} showed several pericentromeric and centromeric signals, chromosome 4H^{ch}S and 5H^{ch}L showed 214 several interstitial signals and chromosome 2H^{ch}L showed a strong interstitial signal. Chromosomes 1H^{ch}L 215 and 3H^{ch}L showed a strong terminal signal. Chromosomes 1H^{ch}S and 4H^{ch}S showed occasionally a weak 216 telomeric signal, and 2H^{ch}L and 7H^{ch}L a weak interstitial one (Fig. 3). Hordeum chilense H7 showed less 217 signals than H1 and H16. Chromosomes 2H^{ch}, 3H^{ch}, 4H^{ch}, 5H^{ch} and 6H^{ch} showed several pericentromeric 218 signals, and chromosomes 4H^{ch}S and 7H^{ch}S showed several interstitial signals. Chromosome 2H^{ch}L showed 219 a strong interstitial signal. Chromosomes 1H^{ch}L, 3H^{ch}L and 7H^{ch}S showed a terminal signal (Fig. 3). The 220 hybridization pattern detected with the GAA probe on H. vulgare was quite different to the pattern observed

in *H. chilense*. All chromosomes showed centromeric or pericentromeric signals, with only chromosome
3H^vL showing a strong distal signal (Fig. 3). As mentioned above, in wheat, the GAA probe hybridized to
all B-genome chromosomes, with strong signals distributed along the whole chromosomes (Fig. 5). All
chromosomes from the A-genome plus 1DS, 7DS and both arms of 2D, also showed some GAA signal, but
the number and intensity were much lower than the ones observed in the B-genome (Fig. 5). The GAA
signals pattern agreed with the one described by Pedersen and Langridge (1997).

227 The (AAC)₅ probe produced an intense signal at the centromeric region of all H1, H16 and H7 228 chromosomes. An exception was chromosomes 5H^{ch}, which did not show any signal, and 7H^{ch}, which only 229 showed an interstitial signal on the long arm (Fig. 3). This probe was quite conserved in the three H. chilense 230 accessions used in this work. The only difference was on chromosome 4H^{ch}L, which showed some 231 interstitial signals in H1 and H16 accessions, but not in H7. In H. vulgare, the (AAC)₅ probe hybridized to 232 the centromeric regions of all chromosomes (Fig. 3). In wheat, this probe hybridized to all B-genome 233 chromosomes, and it is mainly distributed around the centromeric region, although some interstitial signals 234 were also observed in some of the chromosomes (Fig. 5). Chromosomes 2AS, 4AL and 7AL also showed 235 (AAC)₅ signals close to the centromere (Fig. 5).

236 The hybridization pattern of (CTA)₅ was similar in H1 and H16, hybridizing both to the centromeric 237 regions of chromosomes 2H^{ch} and 3H^{ch}, and to 4H^{ch}S in the case of H16 (Fig. 3). However, H7 showed a 238 higher number of signals, with all chromosomes except for 5H^{ch} showing signal (Fig. 3). In H7, signals were pericentromeric on chromosomes 1H^{ch}L, 2H^{ch}L, 3H^{ch}L and 7H^{ch}L and on both arms of 4H^{ch}. On 239 240 chromosome 6H^{ch}, the signal was located at the NOR region (Fig. 3). Hordeum vulgare showed (CTA)₅ 241 signals on all chromosomes, except for chromosomes 1H^v and 3H^v. It hybridized to the pericentromeric and 242 subtelomeric regions on chromosomes 5H^vL and 6H^vS and only to the subtelomeric region on chromosomes 243 2H^vL, 4H^vL and 7H^vL (Fig. 3). In wheat, the (CTA)₅ probe hybridized interstitially to chromosomes 2AL, 244 3AL, 5AL, 7AL and 7BL, and at the centromeric region on 2B and 3B (Fig. 5).

The 4P6 probe did not produce any signal in any of the *Hordeum* used in this study (Fig. S1). This probe only hybridized to the wheat D-genome (Fig. 5). In wheat, signals were detected on five D-genome chromosome pairs: on chromosomes 2DL, 4DL, 5DS, 6DS, and on both arms of chromosome 1D (Fig. 5).

The (AG)₁₂ probe, as 4P6, was absent in all *Hordeum* species used in this work (Fig. S1). In wheat,
some signals were detected on the pericentromeric region of chromosomes 3BS, 5BS and 6BL (Fig. 5).

250

251 Repetitive sequences hybridized to ribosomal DNA: pTa71 and pTa794

- 252 The pTa71 probe is a 9-kb EcoRI fragment of the 18S-25S rDNA isolated from *T. aestivum* (Gerlach
- and Bedbrook 1979). The pTa794 probe is a 410-bp BamHI fragment of the 5S rDNA isolated from T.
- aestivum (Gerlach and Dyer 1980).

Probe pTa71 did not show any difference among the *Hordeum* genotypes used in this study. Signals were detected on the 2 pairs of chromosomes with nucleolar organizing regions (NOR): 5H^{ch}S and 6H^{ch}S in *H. chilense*, and 5H^vS and 6H^vS in *H. vulgare* (Fig. 4). Our results agreed with the results previously published in numerous cytological studies (Cabrera et al. 1995; Szakács et al. 2013; Delgado et al. 2016).

259 On the contrary, the pTa794 probe showed a different pattern among the *Hordeum* studied. In *H*.

260 chilense H1, H16 and H7, this probe was only detected on chromosome 5H^{ch}S (Fig. 4). However, in H.

261 *vulgare*, signals were detected on chromosomes 2H^vL, 4H^vL and 7H^vL (Fig. 4). In wheat, the pTa71 probe

was detected on chromosomes 1BS, 6BS and 5DS, and the pTa794 probe was observed on chromosomes

263 1AS, 1BS,1DS, 5AS and 5BS (Fig. 5).

264

265 Repetitive sequences hybridized to centromeric regions: BAC7, CRW and CCS1

266 The BAC7 probe is a centromere-specific large insert clone from *H. vulgare* L. (Hudakova et al. 2001).

267 The CRW probe is a wheat centromeric retrotransposon from Ae. speltoides Tausch. and Ae. tauschii Coss.

268 (Liu et al. 2008). The CCS1 probe is a 260 bp region within the clone (Hi-10) isolated from B. sylvaticum

269 L. (Abbo et al. 1995; Aragón-Alcaide et al. 1996).

The centromeric probe BAC7 (Hudakova et al. 2001) was specific to *H. vulgare*, labelling the
centromeres of all chromosomes (Fig. 6). Neither *H. chilense* nor *T. aestivum* showed any signals when this
probe was used (Fig. 6).

The centromeric probe CRW (Liu et al. 2008) was detected on all chromosomes of the three accessions of *H. chilense* (H1, H16, H7) and *H. vulgare* (Fig. 6). However, non-specific signals were also frequently observed along chromosomes (Fig. 6). In wheat, probe CRW also hybridized to the centromeric regions of all chromosomes (Fig. 6). Unlike the genus *Hordeum*, signals on wheat were strong and clear, labelling

exclusively the centromeric region. Chromosomes from the D-genome showed weaker signals than A and
B-genome chromosomes, which is due to the fewer number of CRW copies as previously described (Liu et
al. 2008).

The CCS1 probe (Aragón-Alcaide et al. 1996) was also detected in H1, H16, H7, *H. vulgare* and T. aestivum at the centromeric region (Fig. 6). However, as happened with the CRW probe, non-specific signals were frequently observed along chromosomes (Fig. 6). In wheat, the CCS1 pattern was the same as CRW: signals were strong, labelled the centromeric region and chromosomes from the D-genome showed washer signals than A and B genome chromosomes (Fig. 6).

weaker signals than A and B-genome chromosomes (Fig. 6).

285 Up until now, no *H. chilense* specific centromere probe has been described. Therefore, in an attempt to

286 identify *H. chilense* centromeres in the background of wheat, both probes were tested in the *Tritordeum*

287 line "HT27" (described in Cabo et al. 2014a). All wheat chromosomes showed CRW and CCS1 signals as

288 expected; however, surprisingly, *H. chilense* chromosomes did not show any signal with any of the probes

(Fig. S2). On the other hand, the probe (AAC)₅, which hybridized around the centromeres in these species,

showed signals on *H. chilense* chromosomes in the background of bread wheat (Fig. S2).

291

292 Discussion

293 Triticeae species have large genomes, which are primarily composed of repetitive sequences. Many of 294 these repetitive sequences have been used as probes in FISH analysis for genome differentiation, 295 phylogenetic relationship analysis and chromosome identification among different species of the Triticeae 296 tribe (Cabrera et al. 1995; Taketa et al. 2000; Hagras et al. 2005; Jiang and Gill 2006; Marín et al. 2008; 297 Cuadrado et al. 2013; Komuro et al. 2013). The identification and comparison of H. chilense and H. vulgare 298 chromosomes have been carried out in several studies (Hagras et al. 2005; Szakács et al. 2013), however, 299 only a small number of probes and one accession of *H. chilense* were used. For the comparison of both *H.* 300 chilense and H. vulgare with bread wheat, there is barely any published studies where all individual 301 chromosomes are targeted. In this work, our aim was to provide a clear karyotype of H. chilense, H. vulgare 302 and T. aestivum chromosomes, which could be useful in wheat breeding programs to monitor H. chilense 303 and H. vulgare introgressions into wheat, but also to identify the wheat chromosomes where alien segments 304 have been introgressed (Miller et al. 1982; Atienza et al. 2007; Calderón et al. 2012; Rey et al. 2015a, 305 2015b). Moreover, H. chilense and wheat karyotypes would be useful in the development of tritordeum

306 lines, to monitor the chromosome constitution until stable lines are obtained, and to detect spontaneous

307 reorganizations which can occur between *Hordeum* and wheat (Prieto et al. 2001; Cabo et al. 2014b;

308 Delgado et al. 2016; Pujadas Salvá 2016; Delgado et al. 2017).

309 Here, fourteen repetitive probes were used, which accurately identified all individual chromosomes 310 from three accessions of H. chilense (H1, H7 and H16), H. vulgare and T. aestivum. Briefly, the 4P6 and 311 the (AG₁₂) sequences were specific to wheat, and the BAC7 sequence was specific to *H. vulgare*. None of 312 the probes described here was specific to *H. chilense*. The pSc119.2 probe hybridized to both wheat and *H.* 313 chilense, but not to H. vulgare. The rest of the probes hybridized to all three species. At the individual 314 chromosome level, using the pSc119.2, the HvT01 and the GAA probes together, we could identify each 315 individual chromosome from the three accessions of H. chilense (H1, H16 and H7) and H. vulgare. In bread 316 wheat, by using the pAs1 and the GAA probe together we could identify all individual chromosomes, and 317 then by combining them with GISH to label the wheat genome, it was possible to differentiate every wheat 318 chromosome from the ones from barley (both H. chilense and H. vulgare). The remaining repetitive 319 sequences used in this study add further useful information, which, as indicated before, can be used in 320 phylogenetic relationship analysis and when the identification of small chromosome fragments is required.

321 Three centromeric probes, BAC7, CCS1 and CRW were used in this study. Probe BAC7 was specific 322 to H. vulgare as previously described (Hudakova et al. 2001), so no signal was detected in H. chilense and 323 wheat. In the case of both CCS1 and CRW, they are conserved in the three species, labelling the centromeres 324 of all chromosomes (Fig. 6). However, the hybridization signal on wheat was much stronger and clearer 325 than in barley species (H1, H16, H7 and *H. vulgare*), which frequently showed some background signals. 326 Moreover, a confusing result is that neither the CCS1 nor the CRW probes, showed any signal on H. 327 chilense chromosomes when present in the background of tritordeum (Fig. S2). In tritordeum line HT27, wheat chromosomes always showed a centromeric signal, however, this signal was absent in H. chilense 328 329 chromosomes. A possible explanation for this result is that barley carries fewer copies of both CCS1 and 330 CRW and when placed in the background of wheat, the signal is too weak to be detected. This is observed 331 in the case of the wheat D-genome, which carries less copies of CCS1 and CRW sequences compared to 332 the A and B-genome, and accordingly, the hybridization signal in weaker (Liu et al. 2008). None of the 333 centromeric sequences described, can therefore be used to identify the centromeric region of H. chilense 334 chromosomes in tritordeum lines. However, an alternative is to use the (AAC)₅ probe. This repetitive 335 sequence, hybridizes to the centromeric and pericentromeric region of all chromosomes of H. chilense,

except chromosome 5H^{ch} and 7H^{ch}, and to all chromosomes of *H. vulgare*. We tested the (AAC)₅ sequence
in tritordeum line HT27, and *H. chilense* chromosomes were perfectly labelled (Fig. S2). Therefore, the
(AAC)₅ sequence can be used to identify the centromeric region of most *H. chilense* chromosomes in
tritordeum or any other wheat background.

340 It has been suggested that *H. chilense* consists of at least three morphologically and genetically distinct 341 subspecific taxa or groups (Vaz Patto et al. 2001). Repetitive DNA sequences are the main components of 342 heterochromatin and are subject to rapid change, therefore, changes in the distribution of repetitive DNA 343 sequences can provide information of genome evolution and speciation. In this study, we selected one 344 accession of each of the three groups described, to check whether the variability observed morphologically 345 and with molecular markers, is also confirmed by FISH using the fourteen repetitive sequences. Hordeum 346 chilense H1 belongs to group I, H16 to group II and H7 to group III. The results obtained in this work, 347 support the presence of the three different groups. The centromeric probes and the NOR-associated probes 348 did not show any difference among the different H. chilense accessions as expected, since they are very 349 conserved regions. However, the seven repetitive sequences (pAs1, pTa-535, pSc119.2, HvT01, GAA, 350 (AAC)₅ and (CTA)₅) did differ. Probe (CTA)₅ showed the same hybridization pattern in H1 and H16, which 351 differed from the one of H7. In the case of pAs1, pTa-535, pSc119.2, GAA and (CTA)₅, the hybridization 352 pattern was different in the three H. chilense accessions. Based on the FISH results obtained here, accessions 353 H1 and H16 share more similarity between them than with H7. Although these results indicate that the 354 selected set of probes could be useful for a phylogenetic analysis of the different groups described in H. 355 chilense, these three accessions are only one of the many accessions included in each of the groups 356 described, and to confirm the existence of different hybridization pattern in each group, more individuals from each group need to be analyzed. 357

Using the pAs1 probe, Cabrera et al. (1995) suggested that the D-genome from wheat, was the closest phylogenetically to the *H. chilense* genome. We were hoping to add more information supporting this result by using the fourteen probes described in this work. However, only the probes pAs1 and pTa-535, (which show a very similar hybridization pattern to pAs1) have shown some similarity between the D and H^{ch} genomes.

363 The hybridization pattern obtained in this work agreed with what it has been published before except 364 for the (AG_{12}) sequence. We identified three pairs of wheat chromosomes (3BS, 5BS and 6BL) using this 365 probe, while Cuadrado et al. (2008) also describes a signal on chromosome 4B. This difference in the result 366 is probably a consequence of the different FISH conditions used in each study. At this point, it is worth 367 mentioning that some FISH signals can be altered by several factors such as the chromosome spread, the 368 quality of the probe or even the sort of microscope used in the study. Moreover, the examination and 369 identification of the hybridization pattern obtained using repetitive sequences is a complex process that 370 requires experience and previous knowledge of the chromosomal morphology of the species studied. The 371 chromosome spread is critical for obtaining good hybridization patterns when individual chromosomes, 372 chromosome arms or smaller chromosome regions are identified. Some of the hybridization patterns 373 described in this work have already been described previously using different methods for chromosomes 374 spread preparations such as the use of colchicine, ice cold water or nitrous oxide gas (N_2O) (Cabrera et al. 375 1995; Taketa et al. 1999, 2000; Komuro et al. 2013; Tang et al. 2016). These three treatments are 376 extensively used in cytogenetic analysis. However, in our experience, N₂O treatment is the quickest, most 377 reliable and most reproducible method. Here, we combined the protocols from Cabrera et al. (2002) and 378 King et al. (2017) and provide a detailed protocol of how all the FISH experiments were performed in this 379 study.

In summary, we use fourteen repetitive probes to create several karyotypes of *H. chilense* (accessions H1, H16, H7), *H. vulgare* and *T. aestivum;* which together, allow for the identification of every single chromosome in all three species. Moreover, we identify large polymorphism in the three accessions of *H. chilense* studied, which supports the proposal of the existence of different groups inside *H. chilense* species.

384

385 Author contributions

These authors made the following contributions to the manuscript: M-D.R. and A.C.M. designed the research and wrote the manuscript. M-D.R. performed the research and analyzed the data. All authors read and approved the final manuscript.

389

390 Compliance with ethical standards

391 Conflict of interest The authors declare that they have no conflict of interest.

392

393 Research involving human participants and/or animals

- 394 No research involving human participants or animals was performed.
- 395

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- 571 Tables
- 572 Table 1. List of repetitive sequences with their primer sequences, annealing temperatures and
 573 polymerase enzymes used in PCR for the development of FISH probes. F and R mean forward and
 574 reverse primers, respectively.
- 575
- 576 Figure legends

- 577 Fig. 1. FISH patterns (a) and signal distributions (b) of pAs1 and pTa535 on mitotic metaphase
- 578 chromosomes of *H. chilense* (H1, H16 and H7) and *H. vulgare*.
- 579 Fig. 2. FISH patterns (a) and signal distributions (b) of pSc119.2 and HvT01 on mitotic metaphase
- 580 chromosomes of *H. chilense* (H1, H16 and H7) and *H. vulgare*.
- 581 Fig. 3. FISH patterns (a) and signal distributions (b) of GAA, (AAC)₅ and (CTA)₅ on mitotic
- 582 metaphase chromosomes of *H. chilense* (H1, H16 and H7) and *H. vulgare*.
- 583 Fig. 4. FISH patterns (a) and signal distributions (b) of pTa71 and pTa794 on mitotic metaphase
- 584 chromosomes of *H. chilense* (H1, H16 and H7) and *H. vulgare*.
- 585 Fig. 5. FISH patterns (a) and signal distributions (b) of the fourteen repetitive sequences used in this
- 586 study on mitotic metaphase chromosomes of *T. aestivum* cv. Chinese Spring.
- 587 Fig. 6. Amplification by PCR and FISH patterns of BAC7, CCS1 and CRW of *H. chilense* (H1, H16
- 588 and H7), H. vulgare and T. aestivum cv. Chinese Spring. The PCR products were visualized on 2 %
- agarose gels stained with ethidium bromide. L means 100-bp ladder as size marker (Solis BioDyne, Tartu,
- 590 Estonia).
- 591
- 592 Supporting Information
- 593 Fig. S1. Absence of FISH signal using 4P6 and (AG)₁₂ probes on mitotic metaphase chromosomes of
- 594 *H. chilense* (H1, H16 and H7) and *H. vulgare*.
- 595 Fig. S2. FISH pattern of CCS1, CRW and (AAC)₅ probes and identification by GISH of all *H. chilense*
- 596 chromosomes on mitotic metaphase chromosomes of tritordeum "HT27".











