Aligning a New Reference Genetic Map of *Lupinus angustifolius* with the Genome Sequence of the Model Legume, *Lotus japonicus*

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Abstract

We have developed a dense reference genetic map of *Lupinus angustifolius* (2n = 40) based on a set of 106 publicly available recombinant inbred lines derived from a cross between domesticated and wild parental lines. The map comprised 1090 loci in 20 linkage groups and three small clusters, drawing together data from several previous mapping publications plus almost 200 new markers, of which 63 were gene-based markers. A total of 171 mainly gene-based, sequence-tagged site loci served as bridging points for comparing the *Lu. angustifolius* genome with the genome sequence of the model legume, *Lotus japonicus* via BLASTn homology searching. Comparative analysis indicated that the genomes of *Lu. angustifolius* and *Lo. japonicus* are highly diverged structurally but with significant regions of conserved synteny including the region of the *Lu. angustifolius* genome containing the pod-shatter resistance gene, *lentus*. We discuss the potential of synteny analysis for identifying candidate genes for domestication traits in *Lu. angustifolius* and in improving our understanding of Fabaceae genome evolution.

Key words: synteny; genome evolution; narrow-leafed lupin; Fabaceae

1. Introduction

Comparative genetics and genomics use knowledge from different species to define important evolutionary relationships, an important aspect of which is the physical and genetic synteny (two or more homologous loci found on a single chromosome) and collinearity (conserved linear order of loci) between any two species that can be used to explain chromosomal rearrangements. Synteny and collinearity have been widely studied in legumes (Fabaceae family), and these studies have shown direct correspondences between evolutionary distance and the degree of synteny and collinearity. These include whole chromosome levels of synteny between members of the closely related Vicieae1,2 with successively more fractionated and rearranged...
chromosomal relationships moving out to the Loteae, Cicereae and Phaseoleae.

The *Lupinus* genus—part of the Genistoid clade of the Papilionoid legumes—includes a number of agricultural crop species, including *Lupinus angustifolius* L. (narrow-leafed lupin). The Genistoid clade is thought to be one of the first clades to diverge from the rest of the Papilionoid legumes about 56.4 million years ago and is quite distinct from the other clades containing crop or pasture species. The *Lupinus* genus therefore represents a useful outgroup for understanding genome evolution within the legume family.

*Lupinus angustifolius* is diploid (2n = 40 chromosomes) with a nuclear genome size of 2C = 1.89 pg. Cygenetic and linkage analyses found evidence of ancient polyploidy in *Lupinus angustifolius*, but it is functionally diploid. Alignment of a low-density genetic map of *Lupinus angustifolius* to an early draft genome sequence of the model legume species *Medicago truncatula* revealed short regions of conserved synteny between these highly divergent legume species.

Genetic map resources for *Lupinus angustifolius* are modest, comprising two incomplete maps, several gene-tagging reports and physical descriptions of the chromosomes by cytogenetic approaches. The purpose of this current study was to bring together marker data from previous mapping studies, add almost 200 new markers and develop a high-density consensus reference genetic map for *Lupinus angustifolius* with all chromosomes richly populated with gene-based markers. We then used this map to compare the genome structure of *Lupinus angustifolius* with the recently released genome sequence of the model legume, *Lotus japonicus*. By this approach, we hoped to gain further insight into the evolution of Papilionoid legumes by comparing the genomes of a Genistoid species and a Loteae species. We also sought to develop a resource for map alignment-based identification of candidate genes from *Lotus japonicus* for use in *Lupinus angustifolius* genetic studies.

2. Materials and methods

2.1. Characteristics of the *Lupinus angustifolius* mapping population

The *Lupinus angustifolius* recombinant inbred line (RIL) mapping population used in this study was developed at the Department of Agriculture and Food Western Australia (Perth, Australia) and comprised 106 RILs derived from a cross between a domesticated line (breeding line 83A:476) and a wild landrace from Morocco (P27255). These two crossing parents differed in six major genes for key domestication traits used in all current Australian cultivars: *Ku* (early flowering); *iucundis* (low seed alkaloid); *tardus* and *lentus* (pod shatter resistance); *mollis* (water permeable seed) and *leucospernum* (pigmented flowers, seeds and cotyledons, used as a visible marker to indicate when undesirable cross-pollination to wild types has occurred). Two previous mapping studies used subsets of this population: 89 RILs were used by Boersma et al. and 93 RILs were used by Nelson et al. These studies shared 76 RILs in common, which allowed the straightforward combining of data sets from these previous studies.

2.2. Previously published markers

Genotyping data for 522 microsatellite-anchored fragment length polymorphism (MFLP) loci and five domestication genes (*Ku*, *iucundis*, *lentus*, *mollis* and *leucospernum*) were previously reported by Boersma et al. The anthracnose resistance gene and linked marker (*Lanr1* and *AntjM2*, respectively) were reported by Yang et al. and You et al. Genotyping at 298 mainly gene-based sequence tagged site (STS) loci, 74 amplified fragment length polymorphism (AFLP) loci and five domestication genes (*Ku*, *iucundis*, *lentus*, *mollis* and *leucospernum*) was reported in Nelson et al. A sixth domestication gene (*Tardus*), along with three linked markers (TaM1, TaM2 and TaM3), was reported by Boersma et al. Four further markers tagging domestication traits (MoA, LeM1, LeM2 and KuHM1) were reported by Boersma et al. Six isozyme markers were used to screen the population using the methods of Wolko and Weeden.

2.3. New PCR-based STS markers

Forty-two STS primer pairs were developed within the framework of the 6th EU FP Grain Legumes Integrated Project (GLIP) and were provided by the Laboratory of Plant Genetics and Breeding at the Agricultural Biotechnology Center, Godollo, Hungary. The primer pairs were designed on the basis of information on *Medicago truncatula* and *Pisum sativum* consensus sequences using an intron-targeted strategy to amplify single or low copy genes. The primer information is available on the following website: http://bioweb.abc.hu/cgi-rt/mt/pisprim/pisprim.pl.

Nineteen previously unmapped *Lupinus*-derived STS primer pairs based on polymorphism screening conducted in previous comparative mapping studies were used to genotype the RIL population. Two cross-legume primer pairs (PPE and SGR) were provided by Prof. Norman Weeden (Department of Plant Sciences and Plant Pathology at Montana State University, Bozeman, MT, USA). The design strategy was similar to that used for the GLIP marker generation (N. Weeden, pers. comm.).
One BAC-end sequence tag (BEST) marker was developed from the clone 15L10 from the BAC library of \textit{Lu. angustifolius} reported by Kasprzak \textit{et al.}.\textsuperscript{20} BAC DNA was isolated by QIAprep Spin Miniprep Kit (Qiagen, Doncaster, VIC, Australia) and the insert ends were sequenced by BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Scoresby, VIC, Australia) using an AB PRISM 3130xl Genetic Analyzer. On the basis of the BAC-end sequence, primer pairs were designed with aid of Primer3 software (http://frodo.wi.mit.edu/primer3).

An improved dCAPS assay (d212Len) was developed to replace the CAPS marker 212Len reported by Nelson \textit{et al.}\textsuperscript{7} STS primer pairs from the above sources were screened on \textit{Lu. angustifolius} parental DNA. Primer pairs showing large length polymorphisms were used directly to genotype the RIL population. Amplicons of primer pairs showing no visible length polymorphisms were purified and directly sequenced. DNA polymorphisms were identified by manual inspection of alignments and chromatograms and suitable SNaPshot (AB), CAPS (cleaved amplified polymorphic sequence) or dCAPS (derived-CAPS) assays were developed to genotype the RIL population.

2.4. New MFLP markers

An additional 134 MFLP markers were developed using the approach described by Boersma \textit{et al.}.\textsuperscript{9} Briefly, DNA from each RIL was digested by the restriction enzyme \textit{Tru}91 (Roche Diagnostics Australia Pty Ltd, Kew, Australia), an isoschizomer of \textit{MseI}. An AFLP \textit{MseI}-adaptor\textsuperscript{21} was ligated to the restriction fragments using T4 DNA-ligase (Roche). The DNA was then digested a second time with \textit{HpaII} (Gene Works Pty Ltd, Australia) as described by Yang \textit{et al.}.\textsuperscript{22} Pre-selective amplification of fragments using the simple sequence repeat primers listed in Supplementary Table S1 in combination with an \textit{MseI} primer with one selective nucleotide C was followed by a second round of amplification using 16 \textit{MseI} primers having two additional selective nucleotides (Supplementary Table S1). The PCR products were resolved on 5% denaturing sequencing gels and polymorphisms visualized by autoradiography.\textsuperscript{22}

2.5. Linkage map construction

Linkage mapping was conducted with the aid of MultiPoint 1.2 (MultiQTL Ltd, Haifa, Israel), a mapping software package that used the ‘evolutionary optimization strategy’\textsuperscript{23} to resolve locus order. Of the 1118 marker and trait loci inputted to MultiPoint 1.2, 16 marker loci showed severe segregation distortion ($\chi^2 P < 0.001$). These were excluded from further analyses due to their tendency to introduce false linkages into the analysis. Because a large number of loci were included in this analysis, moderately distorted loci ($0.001 < P < 0.01$) were retained but kept under observation in the event that they became implicated in false linkages at a later stage.

Initial clustering into linkage groups was performed at a maximum observable recombination frequency (rf) of 0.10. Multipoint marker ordering was performed separately for each linkage group, and the robustness of marker order was assessed using jackknife re-sampling. Markers ordered with jack-knife values $>90\%$ were considered highly robust and designated as ‘framework’ markers. Markers ordered with jack-knife values $<90\%$ (including redundant markers that mapped to the same location) were initially excluded from the map construction due to their destabilizing effect on locus order. The same set of procedures was carried out serially at gradually increasing maximum recombination frequencies (rf = 0.15, 0.20, 0.25, 0.28, 0.30 and 0.31). At each new clustering cycle, manual inspection of proposed new clusters assisted the identification of valid and invalid clusters. A new cluster was considered valid if its two progenitor clusters were most closely related to each other via their terminal loci. A new cluster was considered invalid if its two progenitor clusters were most closely related by non-terminal loci. Such invalid clusters were associated with problematic loci that showed either high degree of missing data points and/or moderately severe segregation distortion ($0.001 < P < 0.01$) of alleles towards one of the two founding parents of the population. Only clusters judged valid were permitted during linkage mapping.

Once the framework map construction was completed, the interval sizes were transformed to account for multiple meioses involved in the development of the RIL population and were expressed in Kosambi centiMorgans (cM). Markers that had earlier been removed from the map due to their destabilizing effect on locus order (i.e. those with jack-knife values $<90\%$) were assigned to the most likely intervals on the framework map and were referred to as ‘attached’ markers.

2.6. Comparison of \textit{Lu. angustifolius} and \textit{Lo. japonicus} genomes

The \textit{Lo. japonicus} genome sequence version 4 was used to generate pseudomolecules representing the six chromosomes of \textit{Lo. japonicus}. The pseudomolecules were assembled based on the Kazusa clone lists (http://www.kazusa.or.jp/lotus/clonelist.html) using custom Perl scripts (http://www.perl.org/). Comparison of the genetic map of \textit{Lu. angustifolius}
Figure 1. Linkage map of the *Lu. angustifolius* genome comprising 1090 loci distributed among 20 linkage groups (NLL-01 to NLL-20) and three small clusters (Cluster-1 to Cluster-3). Loci on the left of each linkage group are ‘framework’ markers used to construct the linkage group. Loci on the right of each linkage group are ‘attached’ markers whose approximate location between flanking markers is shown. Locus positions are in Kosambi centiMorgans. Major gene trait loci are shown in bold text.
Figure 1. Continued
Figure 1. Continued
and the genome sequence of *Lo. japonicus* was achieved via blastall BLASTn homology search\(^{24,25}\) (at expected alignment values of \(1 \times 10^{-20}\)) and visualized using CMAP hosted at LegumeDB\(^{26}\) and GridMap 3.0 as described by Nelson and Lydiate.\(^{27}\)

### 3. Results and discussion

#### 3.1. A new reference map for *Lu. angustifolius*

The new genetic map of *Lu. angustifolius* comprised 1090 loci arranged in 20 linkage groups and 3 small clusters (Fig. 1; larger scale diagrams are provided in Supplementary Fig. S1). The genotype data and map positions are provided in Supplementary Table S2. The total length of the map was 2361.8 cM, with linkage groups ranging from 69.7 to 168.1 cM. Table 1 gives summary details of each linkage group, including their size, the number of framework and total markers, and the equivalent linkage groups from the two previous maps of *Lu. angustifolius* reported by Boersma *et al*.\(^{9}\) and Nelson *et al*.\(^{7}\) These two previous studies lacked both the critical marker density and sophisticated mapping methodology of the current mapping study, which led to significant differences between the maps. Boersma *et al*.\(^{9}\) reported 21 linkage groups; comparison with the new map revealed that two pairs of the Boersma *et al*.\(^{9}\) linkage groups (LG13 and LG21, and LG17 and LG20) coalesced into two linkage groups in this new map (NLL-08 and NLL-10, respectively) (Table 1). One linkage group (NLL-12) in the new map was entirely absent from the Boersma *et al*.\(^{9}\) map. The Nelson *et al*.\(^{7}\) map had 20 linkage groups (the same number as the current map); comparison with the new map revealed that five linkage groups of Nelson *et al*.\(^{7}\) (LG07, LG19, LG05, LG14 and LG20) were illegitimately joined and were divided among 10 of the new linkage groups (Table 1).

This newly constructed genetic map of *Lu. angustifolius* involved a significantly greater number of markers and a more sophisticated mapping approach (the

<table>
<thead>
<tr>
<th>Linkage group</th>
<th>Length (cM)</th>
<th>Framework loci</th>
<th>Total loci</th>
<th>Linkage group in Boersma <em>et al</em>.(^{9})</th>
<th>Linkage group in Nelson <em>et al</em>.(^{7})</th>
</tr>
</thead>
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<tr>
<td>NLL-01</td>
<td>168.1</td>
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<td>LG07 (top) and LG18</td>
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<td>60</td>
<td>LG6</td>
<td>LG09 (top) and LG15</td>
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<tr>
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<td>65</td>
<td>LG11</td>
<td>LG02</td>
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<td>56</td>
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<td>LG10</td>
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<td>LG3</td>
<td>LG16 and LG14 (top)</td>
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<td>66</td>
<td>LG9</td>
<td>LG04</td>
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<td>55</td>
<td>LG13 and LG21</td>
<td>LG05 (top)</td>
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<td>45</td>
<td>LG18</td>
<td>LG11</td>
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<td>54</td>
<td>LG17 and LG20</td>
<td>LG01, LG19 (bottom) and Triplet-1</td>
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<tr>
<td>NLL-11</td>
<td>112.9</td>
<td>36</td>
<td>78</td>
<td>LG2</td>
<td>LG06 and LG20 (bottom)</td>
</tr>
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<td>NLL-12</td>
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<td>31</td>
<td>Unlinked clusters</td>
<td>LG09</td>
</tr>
<tr>
<td>NLL-13</td>
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<td>25</td>
<td>44</td>
<td>LG16</td>
<td>LG07 (bottom) and Pair-3</td>
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<tr>
<td>NLL-14</td>
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<td>25</td>
<td>36</td>
<td>LG15</td>
<td>LG14 (bottom)</td>
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<td>LG12</td>
<td>LG17 and Triplet-2</td>
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<tr>
<td>NLL-16</td>
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<td>17</td>
<td>34</td>
<td>LG19</td>
<td>LG05 (bottom) and Pair-2</td>
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<tr>
<td>NLL-17</td>
<td>99.7</td>
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<td>49</td>
<td>LG8</td>
<td>LG03</td>
</tr>
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<td>NLL-18</td>
<td>90.8</td>
<td>29</td>
<td>45</td>
<td>LG14</td>
<td>LG12</td>
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<tr>
<td>NLL-19</td>
<td>85.6</td>
<td>22</td>
<td>40</td>
<td>LG7</td>
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<td>Cluster-2</td>
<td>1.8</td>
<td>3</td>
<td>3</td>
<td>Unlinked cluster</td>
<td>—</td>
</tr>
<tr>
<td>Cluster-3</td>
<td>1.4</td>
<td>2</td>
<td>2</td>
<td>Unlinked</td>
<td>Unlinked</td>
</tr>
<tr>
<td>Unlinked</td>
<td>—</td>
<td>—</td>
<td>12</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Total 2361.8 637 1101*

\(^{a}\)Framework markers were used to form the linkage groups. Total loci includes both framework and attached loci (see text for details).
An evolutionary optimization strategy) compared with the two previous maps. Jack-knife re-sampling was highly effective in identifying framework markers that were used to construct stable linkage groups onto which lower quality (and redundant) markers were later attached. This approach is particularly helpful in studies where marker data have been pooled from several sources since the quality of marker genotyping is likely to vary due to differing genotyping technologies and/or technical expertise. These factors led to a greatly improved map with the number of linkage groups equalling the haploid number for this species \((n = 20)\).

This map cannot yet be considered saturated with markers since 18 intervals exceed 15 cM and 1 interval exceeds 20 cM (Fig. 1, Supplementary Fig. S1). However, the marker density is ample for most mapping purposes. Eighteen markers showing significant \((P < 0.01)\) segregation distortion were distributed across nine linkage groups, with the majority \((14)\) being dominant MFLP markers. Therefore, these distorted loci are more likely to have arisen from marker genotyping errors than being an indication of imbalance in the RIL mapping population. The population of RILs used to construct this map is available to the research community on request. With these resources, this map can be considered the new reference genetic map for *Lu. angustifolius*.

### 3.2. Comparing the genomes of *Lu. angustifolius* and *Lo. japonicus*

The new genetic map of *Lu. angustifolius* was compared with the recently released genome sequence of the model legume *Lo. japonicus*. This was achieved by comparing the DNA sequences of 311 STS markers (representing 363 mapped loci in *Lu. angustifolius*) to the genome sequence of *Lo. japonicus* using BLASTn homology searching. At the significance threshold of \(P < 1 \times 10^{-20}\), 159 markers (detecting 187 *Lu. angustifolius* loci) matched one or more locations in the *Lo. japonicus* genome. Seven markers (UWA023, UWA097, UWA158, UWA160, UWA270, UWA300 and LSSR18) representing 16 mapped loci in *Lu. angustifolius* detected repetitive sequences in the *Lo. japonicus* genome and were removed from further analysis. The remaining 152 markers (representing 171 mapped loci in *Lu. angustifolius*) detected one or more correspondences in the *Lo. japonicus* genome. In total, there were 242 correspondences between the two genomes, and these are plotted graphically in Fig. 2 and summarized in Table 2.

#### Table 2. Numbers of *Lu. angustifolius* loci with BLASTn \((P < 1 \times 10^{-20})\) correspondences in six *Lo. japonicus* chromosomes (Lj1–Lj6)

<table>
<thead>
<tr>
<th>Lotus japonicus chromosomes</th>
<th>Primary correspondences</th>
<th>All correspondences^a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lj1</td>
<td>40</td>
<td>60</td>
</tr>
<tr>
<td>Lj2</td>
<td>36</td>
<td>52</td>
</tr>
<tr>
<td>Lj3</td>
<td>31</td>
<td>47</td>
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<td>Lj4</td>
<td>36</td>
<td>44</td>
</tr>
<tr>
<td>Lj5</td>
<td>15</td>
<td>22</td>
</tr>
<tr>
<td>Lj6</td>
<td>13</td>
<td>17</td>
</tr>
<tr>
<td>Total</td>
<td>171</td>
<td>242</td>
</tr>
</tbody>
</table>

^aPrimary, secondary and tertiary BLASTn correspondences.

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*Figure 2.* Global distribution of synteny between 20 linkage groups of *Lu. angustifolius* (NLL-01 to NLL-20) and 6 chromosomes of *Lo. japonicus* (Lj1–Lj6). Loci showing homology between the two genomes at \(P < 1 \times 10^{-20}\) significance threshold are indicated by dots.
Figure 3. Three *Lu. angustifolius* linkage groups (NLL-08, NLL-06 and NLL-12) showing conserved gene order with *Lo. japonicus* chromosomes Lj2, Lj3, Lj4 and Lj5. The pod shatter resistance gene, *Lentus*, is located on NLL-08. Names of loci showing conserved synteny between the genomes are shown next to the *Lu. angustifolius* linkage groups (scaled in Kosambi centiMorgans, cM). The start position of homologous sequences on *Lo. japonicus* pseudomolecules is shown next to *Lo. japonicus* chromosomes (scaled in megabases, Mb).
Of the 120 pairwise chromosome comparisons (i.e. 20 \textit{Lu. angustifolius} \times 6 \textit{Lo. japonicus} chromosomes), 50 had at least two correspondences (the minimum for detecting syntenic) and 34 had at least three correspondences (the minimum for detecting collinearity). There were clear differences in the degree of conserved collinearity on \textit{Lu. angustifolius} and \textit{Lo. japonicus} chromosomes with some regions showing good conservation whereas other regions were highly rearranged with respect to each other (Fig. 2). This heterogeneous pattern of genome collinearity was also observed in an earlier genome comparison between \textit{Lu. angustifolius} (a Genistoid species) and \textit{M. truncatula} (a Galeoid species), and provides additional support to the widespread evolutionary distance believed to separate the Galegoids and Genistoids. A similarly heterogeneous pattern was observed between \textit{Arachis hypogea} (groundnut, in the Dalbergioid clade, which is also considered relatively basal in the evolution of Papilionoid legumes) and the Galeoid species \textit{Lo. japonicus} and \textit{M. truncatula}. It is interesting to note that arguably the best conserved region between the \textit{Arachis} and \textit{Lo. japonicus} genomes on chromosomes Ar6 and Lj1 (synteny block 5, delineated by contigs CM0222–CM0105), respectively, appears to show no conservation in the \textit{Lu. angustifolius} genome. A more thorough analysis will be required to make firm conclusions, but on first inspection it would appear that regions of conserved gene order may not be generalized across wide evolutionary distance within the Papilionoid legumes. This is in contrast to the crucifer family where large variations in chromosome numbers and frequent rearrangements exist, but where the integrity of many chromosomal blocks has been maintained over a similarly long evolutionary timeframe.

Three regions showing relatively good conservation of locus order between \textit{Lu. angustifolius} and \textit{Lo. japonicus} are shown in Fig. 3. Even these relatively well-conserved regions show evidence of chromosome translocations and/or inversions, which will make exploitation of syntenic by lupin researchers more difficult. One of the potential applications of syntenic is map alignment-based identification of candidate genes for important traits in the crop species by searching for likely candidates in the equivalent region of the model genome. For example, the pod shatter resistance gene \textit{lentus} is located on \textit{Lu. angustifolius} linkage group NLL-08 in a region with relatively well-conserved gene order with \textit{Lo. japonicus} chromosome 5 (Fig. 3). Unfortunately, there appears to be an inversion in the interval containing \textit{lentus} such that the equivalent region of \textit{Lo. japonicus} is somewhat ambiguous and is only delimited to the lower half of chromosome 5. Clearly, more bridging points will be required to make such map-based inferences in \textit{Lu. angustifolius} using \textit{Lo. japonicus} as a model genome and therefore more gene-based markers are required in \textit{Lu. angustifolius}.

In conclusion, we present a high-resolution reference map for \textit{Lu. angustifolius} which will serve as a shared resource for the legume genetic community. We discovered that the \textit{Lu. angustifolius} genome has numerous chromosomal rearrangements relative to the \textit{Lo. japonicus} genome, though widespread but small sections of conserved gene order are present. As an outlying species compared with other legume model and crop species, \textit{Lu. angustifolius} serves as a useful reference for gaining better understanding of legume genome evolution.

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\textbf{Supplementary data:} Supplementary data are available at www.dnaresearch.oxfordjournals.org.

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