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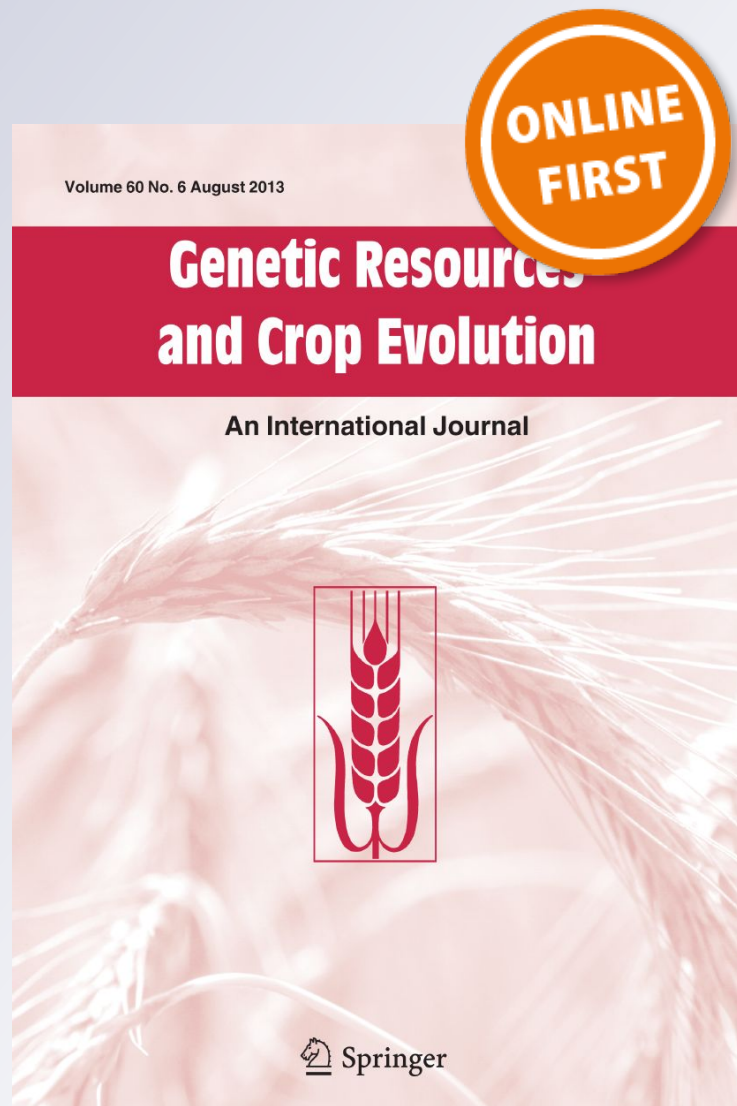
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## RESEARCH ARTICLE

# Persistence of rye (*Secale cereale* L.) chromosome arm 1RS in wheat (*Triticum aestivum* L.) breeding programs of the Great Plains of North America

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**Abstract** Rye (*Secale cereale* L.) chromosome arm 1RS has been used world-wide by wheat (*Triticum aestivum* L.) breeding programs as a source of pest- and pathogen-resistance genes, and to improve grain yield and stress tolerance. The most common vehicles used to access 1RS are various 1AL.1RS and 1BL.1RS wheat-rye chromosomal translocations. Over the past 25 years, advanced North American wheat breeding lines were evaluated, first by assay of secalin storage proteins, and later by use of DNA marker TSM0120, for the presence of these two translocations. Both methods provide accurate and efficient means of identifying and differentiating 1BL.1RS and 1A.1RS. Both 1AL.1RS and 1BL.1RS wheats were found in all tested years. 1AL.1RS lines were more common in southern Great Plains breeding programs. 1AL.1RS lines were released as cultivars at a frequency identical

to that of wild-type breeding lines. In contrast, 1BL.1RS breeding lines were developed by breeding programs throughout the Great Plains, but fewer were released as cultivars. Both 1RS translocation types persist in Great Plains breeding programs. The lower rate of release of 1BL.1RS cultivars no doubt is a consequence of the more drastic effects on breadmaking quality relative to those observed with 1AL.1RS.

**Keywords** Wheat · Rye · Chromosomal translocations · North American breeding programs · DNA protein markers

## Introduction

Common wheat (*Triticum aestivum* L.), a product of interspecific hybridization and chromosome doubling (Kihara 1975; Matsuoka 2011) tolerates the loss or addition of its own chromosomes and those of other members of the Triticeae. In attempts to access potentially useful traits from many closely related species, wheat geneticists introduced alien chromosomes and chromosome fragments into wheat via interspecific hybridization (McFadden and Sears 1947). Some of the earliest successful interspecific hybrids were between wheat and rye (*Secale cereale* L.) (Leighty and Sando 1928; Florell 1931). Modern wheat production has heavily utilized genes from rye, particularly those found on chromosome arm 1RS.

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Robert A. Graybosch: Retired.

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Useful traits found on that arm include resistance to various fungal and viral pathogens and insect pests, more efficient mineral absorption, enhanced stress tolerance, and fertility restoration in hybrid wheat production (Lukaszewski 2015). Rye chromosome arm 1RS is most commonly found in wheat cultivars in the form of 1BL.1RS wheat-rye chromosomal translocations. World-wide, hundreds of 1BL.1RS wheat cultivars have been bred and released (Lukaszewski 2015). In the Great Plains of North America, but less so in other parts of the world, 1AL.1RS cultivars, typically derived from the wheat germplasm line Amigo (Sebesta et al. 1995), are also common.

In 1BL.1RS and 1AL.1RS chromosomal translocations, the short arm of 1R replaces the short arms of either 1B or 1A of wheat. Given that replacement, wheat genes encoding low-molecular-weight glutenin (LMW) and gliadin grain-storage proteins are lost, and rye genes encoding secalin storage proteins are introduced to wheat. In wheats lacking 1RS, LMW glutenin subunits polymerize and add to the glutenin matrix responsible for dough strength. Due to either the loss of LMW glutenins and gliadins, the presence of the more hydrophilic secalins (Hussain and Lukow 1994), or both, processing-quality defects long have been associated with 1RS in wheat backgrounds (Zeller and Hsam 1983; Graybosch 2001). The effect, however, is not as pronounced in 1AL.1RS wheats as in those lines carrying 1BL.1RS (Graybosch et al. 1993).

The first 1RS cultivars released in the Great Plains of North America were the 1BL.1RS cultivar 'Siouxland', released in 1985 by the University of Nebraska (Schmidt et al. 1985) and the 1AL.1RS cultivar 'TAM-107' (Porter et al. 1987), released from Texas A&M University. 'Siouxland' carries the 1BL.1RS translocation derived from the Russian wheat cultivar 'Kavkaz', and 'TAM-107' carries 1AL.1RS derived from wheat germplasm line Amigo (Graybosch 2001). Although both 'Siouxland' and 'TAM-107' were widely grown (Baenziger et al. 1989; Raeburn 1996), and heavily used as parents in North American breeding programs (Graybosch 2001), the release of 'Siouxland' was met with dismay by the wheat milling and baking industries, due to its poor dough strength and baking properties (Martin and Stewart 1990; Graybosch 2001). In the early 1990s, USDA Agricultural Research Service (ARS) scientists at Lincoln, Nebraska began screening advanced US breeding lines

for the presence of 1AL.1RS and 1BL.1RS. This was done to assist breeders in making both release decisions and in selecting parental materials for new matings.

Breeding lines screened by the USDA-ARS primarily were derived from the USDA-coordinated regional nursery trials. Since 1931, the Uniform Regional Nursery Program has tested advanced breeding lines from public and private breeding programs in the US, with entries occasionally also supplied from Canada. Two such nurseries, the Southern (SRPN) and Northern (NRPN) Regional Performance Nurseries (USDA 2018), operate in the Great Plains region of North America. In the SRPN and NRPN, advanced breeding lines are tested in replicated field trials from Texas north to Alberta, Canada, and from the Missouri River west to the Rocky Mountains. The SRPN is tested from southern Texas to South Dakota, with most entries being derived from breeding programs based in Nebraska south to Texas. The NRPN is tested from Kansas north to Alberta, with entries typically from breeding programs based in Nebraska, the Dakotas, and Montana. Lines also are tested for disease and pest resistance, bread-making quality, presence of key molecular markers, and other traits. Lines entered in the SRPN and NRPN typically have already been tested for multiple years in their states of origin, with the regional nursery trials representing one of the last stages before cultivar-release decisions are made. Thus, entries represent the most elite lines from the submitting breeding programs at any given time.

From 1993 to 2010, 1RS was identified in SRPN and NRPN entries via the presence of rye secalin proteins in grain samples, by using sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) separation of grain proteins (Graybosch et al. 1999). Commencing in 2010, USDA-ARS scientists at the USDA Central Small Grain Genotyping Lab (CSGGL) in Manhattan, KS (<https://hwwgenotyping.ksu.edu>) started screening the SRPN and NRPN entries with DNA markers for key traits and genes. In 2010 and 2011, both SDS-PAGE and DNA markers were applied in the 1RS screen. As these initial results showed 100% agreement, the SDS-PAGE work was suspended. Recently, however, we decided to re-visit this topic and screen the 2012–2018 SRPN and NRPN again with SDS-PAGE. This was undertaken to determine the accuracy of the DNA-based markers,

and to verify that IRS was inherited as one large linkage block. Typically, recombination does not occur between IRS and any wheat chromosome arms, unless present in a *Ph*-background (Lukaszewski 2000). Disagreement of SDS-PAGE and DNA-based results could indicate the generation of a rare recombinant that separated secalin-encoding sequences from other genes on IRS. As IRS data now are available on the most advanced wheat breeding lines from Great Plains' programs, a secondary objective was to summarize the past 25 years of results, documenting the impact of IRS on Great Plains wheat breeding and production.

## Materials and methods

### SDS-PAGE

Seed from all SRPN and NRPN 2012–2018 entries was obtained from remnant samples of the original entries maintained in cold storage at the Stewart Seed Lab, University of Nebraska, Lincoln. Eight seeds of each entry were placed in 5 ml screw-top vials along with three 6.35 mm stainless-steel bearing balls per sample and ground to meal with a Geno/Grinder® (SPEX SamplePrep Metuchen, NJ, USA). Samples were ground at 1500 rpm for three min to obtain homogeneous meal. A 25 mg aliquot was weighed into a 1.5 ml conical Eppendorf tube, 1 ml H<sub>2</sub>O added, and samples placed on a laboratory shaker at 150 rpm for 30–60 min. Following centrifugation at 14,000 rpm for 5 min, 300 µl supernatant was removed and added to a fresh tube containing 700 µl 100% ethanol. The solution was stored at – 20 °C for 1 h, recentrifuged, and 200 µl supernatant dried in a Labconco Centrivap SpeedVac Concentrator (Kansas City, MO). Dried samples were re-suspended in 250 µl of sample buffer (Graybosch and Morris 1990) and 5 µl aliquots loaded and separated on 12% Bio Rad Criterion TGX Stain-Free Protein Gels (Heracles, CA), by using a Tris-glycine buffer (Graybosch and Morris 1990). Proteins were visualized with silver-nitrate staining (Graybosch and Morris 1990). Each gel also contained control samples of the cultivars 'Siouxland' (1BL.1RS) and 'TAM-107' (1AL.1RS). Secalins were identified as per Graybosch et al. (1999).

### Identification of secalins by mass-spectrometry

Mass spectrometry of peptides released from protease-digested gel bands was used to confirm the identify of diagnostic proteins for 1BL.1RS and 1AL.1RS. For these experiments, proteins were extracted as described above, except that after the – 20 °C treatment and centrifugation, the entire 1 ml supernatant was dried in the Labconco Centrivap SpeedVac Concentrator. The dried sample was again suspended in 250 µl extraction buffer (above) and 15 µl aliquots loaded on 12% gels as described above. Proteins were visualized after staining with Coomassie Brilliant Blue R-250 (300 mg in 45% v/v methanol, 10% acetic acid, 45% H<sub>2</sub>O). After de-staining with 10% methanol/7% acetic acid, diagnostic bands (see Results and Discussion) were cut from the gels and analyzed at the University of Nebraska Lincoln, Proteomics and Metabolomics Core Facility (<https://biotech.unl.edu/proteomics-and-metabolomics>).

Excised gel bands were washed with ammonium bicarbonate/acetonitrile to remove the stain and SDS. Trypsin was added and digestion carried out overnight at 37 °C, followed by chymotrypsin digestion at 30 °C overnight. Peptides were extracted from the gel pieces, dried down, and re-dissolved in an aqueous solution of 2.5% v/v acetonitrile, 0.1% formic acid. Each digest was separated by nano-LC-MS/MS using a gradient on a 0.075 mm × 250 mm C18 column feeding for 1 h into a Q-Exactive HF mass spectrometer (Thermo Fisher Scientific, Lenexa, KS, USA). All MS/MS samples were analyzed with Mascot (Matrix Science, London, UK; version 2.6.1). Mascot was searched with a fragment ion mass tolerance of 0.060 Da and a parent ion tolerance of 10.0 PPM. De-amidated asparagine and glutamine and oxidized methionine were specified in Mascot as variable modifications. Scaffold (version Scaffold\_4.8.4, Proteome Software Inc., Portland, OR) was used to validate MS/MS based peptide and protein identifications. Peptide identifications were accepted if they could be established at greater than 80.0% probability by the Peptide Prophet algorithm (Keller et al. 2002), with Scaffold delta-mass correction. Protein identifications were accepted if they could be established at greater than 99.0% probability and contained at least 2 identified peptides. Protein probabilities were assigned by the Protein Prophet algorithm (Nesvizhskii et al. 2003). Proteins that contained similar peptides and could not



be differentiated based on MS/MS analysis alone were grouped to satisfy the principles of parsimony. Proteins sharing significant peptide evidence were grouped into clusters using Scaffold 4 analysis tool (Proteome Software, Portland, OR, USA: <http://www.proteomesoftware.com/>).

#### Rye SSR marker TSM120

Rye SSR marker SCM9 (Saal and Wricke 1999) has been used by USDA-ARS genotyping labs (<https://maswheat.ucdavis.edu/protocols/Drought/index.htm>) for detection of 1BL.1RS, and 1AL.1RS translocations. However, SCM9 has sometimes produced both false positives and false negatives at the USDA-ARS CSGGL (data not shown). We discovered a marker that is more reliable for detecting wheat-rye translocations by screening 29 rye SSR markers previously mapped by Kofler et al. (2008) to the distal bin of rye chromosome 1. All 29 rye markers were tested on a panel of wheat lines known to have either wheat-rye translocations or no translocation (data not shown). The panel consisted of 44 wheat cultivars and breeding lines, 22 known to carry 1AL.1RS, 10 with 1BL.1RS, and 12 wild-type (non-translocation) lines. SSR marker TSM120 was found to clearly and repeatably distinguish between non-1RS, 1AL.1RS, and 1BL.1RS wheat-rye translocations. No PCR amplification occurred in any of the 12 wheat lines without 1RS translocations. PCR products from 1AL.1RS translocation lines are 358 bp long, while 1BL.1RS translocation lines produce a 345 bp PCR product. Primer sequences for gel-based detection of TSM0120 fragments are: TSM0120F 5'-CCG CCG TCC TCC TCC T-3' and TSM0120R 5'-AGA CGG CAG GCA TGG AT-3'. To use fluorescently labeled primers in a capillary sequencer, primer TSM0120F was replaced with an 18-bp tailed forward primer: TSM0120F18 5'-ACG ACG TTG TAA AAC GAC CCG CCG TCC TCC TCC T-3'. As expected, the resulting fragment sizes increased by 18 bp.

#### DNA marker analysis

All 2010–2018 entries in the NRPN and SRPN were screened with TSM0120. Each 13  $\mu$ l PCR reaction contained 20–80 ng DNA extracted from 4 to 8 seedlings (except with single-seed samples as noted below), 50 nM of M13-tailed forward primer, 100 nM

of reverse primer, 50 nM fluorescence-labeled M13 primer, 200  $\mu$ M of each dNTP, 1.3  $\mu$ l 10X ammonium sulfate PCR buffer, 2.5 mM MgCl<sub>2</sub>, and 1 unit of *Taq* polymerase (various vendors). PCR was performed by following a 56 °C touch-down program (Liu et al. 2008) in a DNA Engine<sup>®</sup> Peltier Thermal Cycler (Bio-Rad Lab, Hercules, CA, USA). Up to four different plates of PCR products labeled with one of four dyes (FAM, VIC, NED, and PET) were pooled into one plate by using a Biomek NX<sup>P</sup> liquid handling system (Beckman Coulter Inc., Brea, CA, USA). The pooled PCR products were analyzed with an ABI Prism 3730 DNA Sequencer (Applied Biosystems, Foster City, CA, USA), and their fragment lengths and amplicon polymorphisms were scored with GeneMarker (SoftGenetics LLC, State College, PA, USA).

#### Scoring and statistical analysis

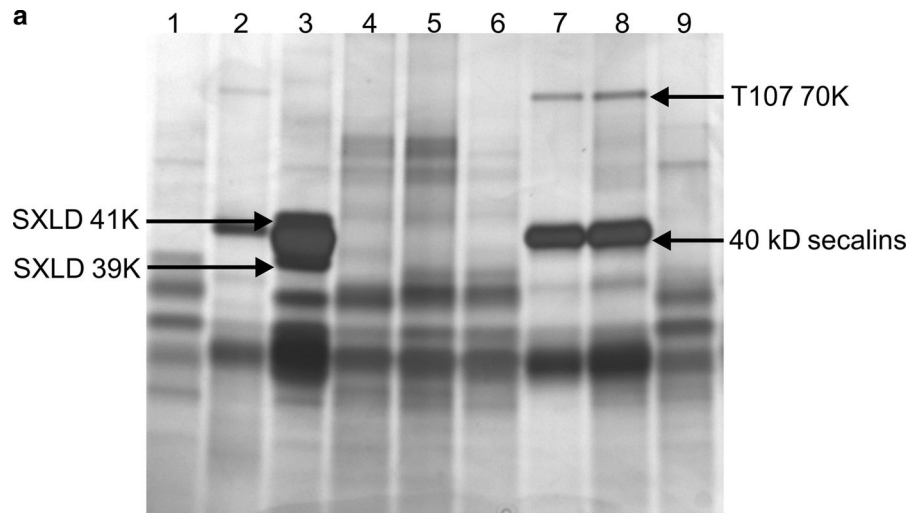
Scoring of samples was double-blind. DNA marker analysis was performed and marker data interpreted, before SDS-PAGE evaluations were conducted. SDS-PAGE analyses were conducted and data scored, without consulting DNA marker results. Samples that disagreed after the first evaluation were retested as single-seed or single-plant samples with both the SDS-PAGE and DNA marker protocols as described above.

The frequency of 1BL.1RS and 1AL.1RS in the SRPN and NRPN was summarized in relation to the number of entries per year. Long-term control cultivars included in the nurseries were excluded from the frequency analysis. No attempt was made to distinguish homogeneous lines from heterogeneous. If 1RS was detected, the sample was scored as such. The number of breeding lines released as cultivars (as of 2018), also was determined for 1BL.1RS, 1AL.1RS, and wild-type (WT) lines. Chi square analysis was used to compare frequencies between the two nurseries, and between released and non-released breeding lines. Computations were performed in Excel.

#### Results and discussion

Identities of the diagnostic proteins used in SDS-PAGE evaluations (Fig. 1a) were validated by analysis of peptides released from tryptic/chymotryptic digests of excised gel bands followed by mass spectrometry. All 1RS lines produce a protein band

**Fig. 1** Enrichment and identification of secalins in IRS wheats. **a** Secalins separated by SDS-PAGE. Bands corresponding to 41, 40 and 39 kD secalins and the unknown 70 kDa protein are marked with arrows. **b** Identification of peptides arising from omega secalins. Peptides identified by mass spectrometry following proteolytic digests of isolated putative secalin bands (see panel **a**) are highlighted in gray



**b**

AAB58403.1 (100%), 41,622.8 Da  
 omega secalin. (*Secale cereale*)  
 4 exclusive unique peptides, 9 exclusive unique spectra,  
 78 total spectra, 179/357 amino acids (50% coverage)

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M K T F L I F V L A M T M S I I T T A R Q L N P S E Q E L Q S P Q Q P V P K E Q
S Y P Q Q P Y P S H Q P F P T P Q Q Y S P Y Q P Q Q P F P Q Q P T P I Q P Q
Q P F P Q Q P Q Q P F S Q P Q Q Q L P L Q P Q Q P F P Q P Q Q P I P Q Q P Q Q S
F P Q Q P Q R P E Q Q F P Q Q P Q Q I I P Q Q T Q Q P F P L Q P Q Q P F P Q Q P
Q R P F A Q Q P E Q I I S Q Q P F P L Q P Q Q P F S Q P Q Q P F P Q Q P E Q I I
P Q Q P Q Q P S P L Q P Q Q P F S Q Q P Q R P Q Q P F P Q Q P Q Q I I P Q Q P Q
Q P F P L Q P Q Q P V P Q Q P Q R P F G Q Q P E Q I I S Q R P Q Q P F L L Q P Q
Q P F S Q P Q Q P F L Q Q P G Q I I P Q Q P Q Q P F P L Q P Q Q P F P Q Q P E Q
I I S E Q P Q Q P F L L Q P Q Q P S P Q Q P Q L P F P Q P Q Q P F V V V V
    
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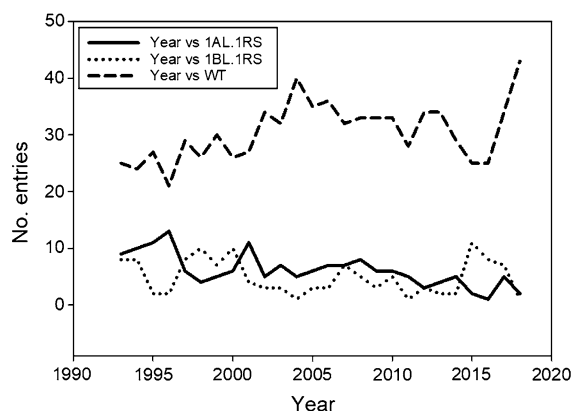
recognized as secalins with an apparent molecular mass of 40 kD (Fig. 1a; Graybosch et al. 1999). In 1BL.1RS lines, two additional bands, herein designated SXLD 41K and SXLD 39K, migrate above and below this large 40 kD band (Fig. 1a). 1AL.1RS lines also produce a unique protein, designated T107 70K. The 40 kD cluster, SXLD 41K, and SXLD 39K all were positively identified as secalins by mass spectrometry (Fig. 1b). T107 70K, however, could not be identified. Peptide sequences of T107 70K were searched against the entire plant database in NCBIprot (<https://www.ncbi.nlm.nih.gov/protein>)—6,481,660 entries—and no matches to secalins (or gliadins) were detected. Protein T107 70K was always found in lines derived from known 1AL.1RS parents and in lines displaying the 1AL.1RS pattern from DNA marker TSM0120. Proteins SXLD 41K and SXLD 39K always co-existed in lines carrying 1BL.1RS as detected by TSM0120.

Exclusive of control cultivars, 525 breeding lines were entered in the 2012–2018 NRPN and SRPN. Results of the double-blind scoring of IRS presence via SDS-PAGE and DNA marker TSM0120 initially agreed on all but nine entries, resulting in an error frequency of 1.7%. DNA marker TSM0120 and SDS-PAGE assays were re-conducted on 12 single seed samples of each of these nine lines, again were scored double-blind. In this second round of screening, five samples were scored non-IRS by both methods, two were scored 1BL.1RS by both methods, and two still disagreed. The two breeding lines in disagreement were scored non-IRS by TSM0120 and heterogeneous by SDS-PAGE. One sample had only one of 12 seeds with secalins, a frequency that could have resulted from mechanical mixing at harvest or human error. The second sample had six of 12 seeds with secalins on SDS-PAGE. DNA marker TSM0120, therefore, is a highly effective rapid screening tool, and the

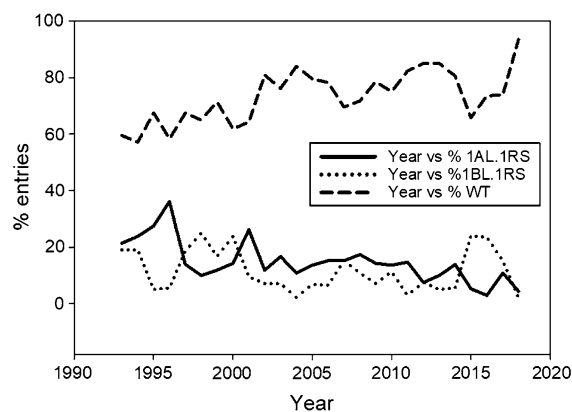
infrequent disagreements between marker and protein data were most likely due to sample heterogeneity, and not due to some rare recombination events. Storage protein and IRS heterogeneity has been reported in Great Plains wheat breeding programs, where selection can commence at the F<sub>3</sub> or F<sub>4</sub> generation (Graybosch 1992; Moreno-Sevilla et al. 1995), whereas recombination between IRS and corresponding wheat chromosomal arms occurs only in the presence of ph- mutants (Lukaszewski 2000).

Results from the 2012–2018 nurseries were combined with those from 1993 to 2011 to examine changes, if any, in frequencies of IRS over time. All lines tested are listed in the Supplementary Materials and also at: <https://www.ars.usda.gov/plains-area/lincoln-ne/wheat-sorghum-and-forage-research/docs/hard-winter-wheat-regional-nursery-program/research/> (USDA 2018). 1AL.1RS wheats were observed at a statistically significant higher frequency (14.7%, Table 1) in the southern Great Plains SRPN than in the northern NRPN (4.3%). 1BL.1RS lines, in contrast, were observed at statistically indistinguishable frequencies (Table 1) in the two nurseries. Within each nursery, the frequency of the two IRS types differed significantly (Table 1) with 1BL.1RS lines being far more common than 1AL.1RS in the NRPN. The difference in the SRPN (Table 1) was slight, but there were significantly more 1AL.1RS lines than 1BL.1RS lines.

Visualization of the frequencies and total numbers of 1AL.1RS and 1BL.1RS in the SRPN and NRPN over time (Figs. 2, 3, 4, 5) shows a relatively constant supply of IRS entries, with some waxing and waning. In the SRPN, 1AL.1RS wheats represented more than 20% of the entries in the early 1990s. This frequency has declined slightly over time, never exceeding 5% in



**Fig. 2** Total number of 1AL.1RS, 1BL.1RS and wild-type (WT) entries, exclusive of controls, Southern Regional Performance Nursery



**Fig. 3** Percentage 1AL.1RS, 1BL.1RS and wild-type (WT) entries, exclusive of controls, Southern Regional Performance Nursery

the most recent 5 years tested. In the NRPN, 1AL.1RS entries rarely exceeded 5% in any of the 25 years evaluated. The frequency of 1BL.1RS in both

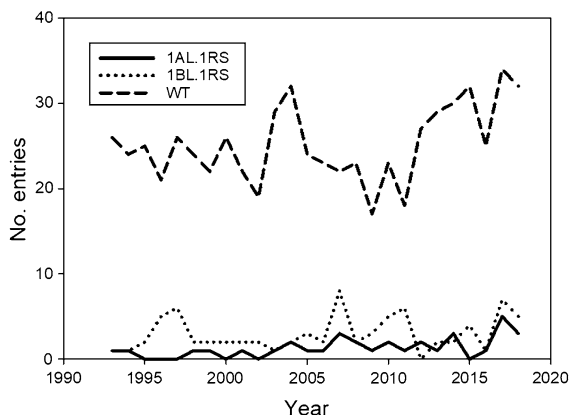
**Table 1** Frequency of IRS wheat-rye translocation lines in USDA-ARS coordinated hard winter wheat breeding trials, 1993–2018

Nursery	Entries tested <sup>a</sup>	1AL.1RS		1BL.1RS		Wild-type		$\chi^2$ (1AL.1RS vs 1BL.1RS)	<i>p</i>
		No.	%	No.	%	No.	%		
NRPN	761	33	4.3	77	10.1	651	88.6	59.4	< 0.001
SRPN	1081	159	14.7	127	11.8	795	73.55	6.4	< 0.025
$\chi^2$ (NRPN vs SPRN)		55.7		1.82					
<i>p</i>		< 0.001		> 0.10					

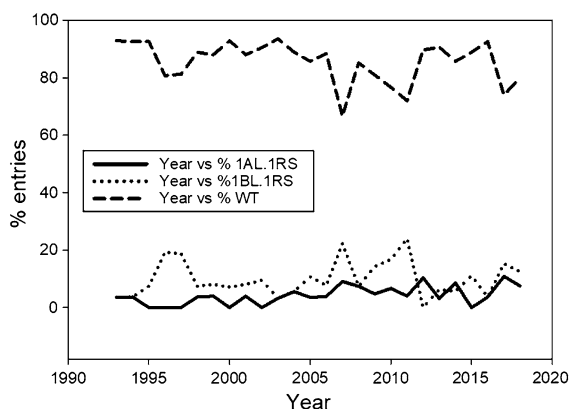
NRPN Northern Regional Performance Nursery, SRPN Southern Regional Performance Nursery

<sup>a</sup>Includes all experimental lines entered; long-term control cultivars removed





**Fig. 4** Total number of 1AL.1RS, 1BL.1RS and wild-type (WT) entries, exclusive of controls, Northern Regional Performance Nursery



**Fig. 5** Percentage 1AL.1RS, 1BL.1RS and wild-type (WT) entries, exclusive of controls, Northern Regional Performance Nursery

nurseries fluctuated. In the SRPN in the late 1990s, 1BL.1RS approached a frequency of 20% for several years, then declined, only to increase markedly around 2015 (Figs. 2, 3). Likewise, 1BL.1RS entries approached or exceeded 15% in the NRPN in the mid-1990s, again in 2006 and 2011 (Figs. 4, 5), and declining during intervals between these years.

Nearly all cultivars released in the Great Plains region are evaluated in either or both the SRPN or NRPN. Of 1040 unique, wild-type breeding lines evaluated in these trials from 1993 to 2018, 180 (as of this writing) have been released as cultivars (Table 2; USDA 2018). The frequency (18.1%) of 1AL.1RS lines released as cultivars, to date, is not significantly different than the frequency (17.3%) of wild-type lines released (Table 2). The frequency (11.4%) of

1BL.1RS breeding lines ultimately released as cultivars, however, is significantly lower than that of wild-type cultivars (Table 2). Great Plains winter wheat 1RS cultivars released from 1993 to 2018 are listed in Table 3.

Observations may be summarized as follows: Both 1AL.1RS and 1BL.1RS wheats have been observed at a relatively constant frequency over the past 25 years in Great Plains breeding programs. 1AL.1RS lines are more common in southern Great Plains breeding programs, which typically enter lines in the SRPN. 1AL.1RS lines are released as cultivars at a frequency identical to that of wild-type breeding lines. In contrast, 1BL.1RS breeding lines are produced by breeding programs across the Great Plains, but fewer are released as cultivars. Both 1RS translocation types persist in Great Plains breeding programs. The lower rate of release of 1BL.1RS cultivars no doubt is a consequence of the more drastic effects on breadmaking quality relative to those observed with 1AL.1RS (Graybosch et al. 1993). Among Great Plains wheat breeders, the perception is that 1AL.1RS has little, if any, negative quality effects. At a recent field day, one such breeder was heard to utter “we don’t fear this translocation” with reference to 1AL.1RS quality.

Recent literature suggests the situation for 1BL.1RS lines is similar in many other parts of the world. 1BL.1RS was observed at frequencies greater than 25% in samples of both elite cultivars and advanced breeding lines from Pakistan tested in 2010 (Tahir et al. 2014). In Bulgaria, 1BL.1RS was observed in 10% of cultivars tested (Landjeva et al. 2006). Thirteen percent of Hungarian wheats tested in 2005 carried 1BL.1RS, although 50% of the cultivars registered in 1994 carried this translocation (Purnhauser et al. 2011). A compendium of released lines with 1RS maintained by Schlegel (2016) lists many hundreds of cultivars world-wide. 1AL.1RS cultivars originally were restricted to the US; however, Lukaszewski (2015) noted that their frequency and distribution outside of the US has increased since the year 2000.

Lukaszewski (2015) further observed “It is interesting that most of the spread of the translocation occurred when, or after, the rye resistance genes have been broken down and contributed little value in breeding.” If most of the resistance genes on 1RS are, in fact, broken, how can one explain its continued presence? Observations suggest that 1RS contributes other

**Table 2** Frequency of cultivar release from breeding lines entered in USDA-ARS coordinated hard winter wheat breeding trials, 1993–2018

Genotype	N <sup>a</sup>	N released as cultivars	% released	$\chi^2$ (vs wild-type)	<i>p</i>
1AL.1RS	138	25	18.1	0.04	> 0.75
1BL.1RS	141	16	11.4	3.24	< 0.10
Wild-type	1040	180	17.3		

<sup>a</sup>N = number of unique genotypes. Redundant entries between SRPN and NRPN omitted

**Table 3** Great Plains 1RS cultivars released 1993–2018

Genotype	Cultivar	Year released	Genotype	Cultivar	Year released
1AL.1RS	Hickok	1993	1BL.1RS	Custer	1994
1AL.1RS	Ogallala	1993	1BL.1RS	Tonkawa	1994
1AL.1RS	Nekota	1994	1BL.1RS	Quantum 7406	1997
1AL.1RS	Niobrara	1994	1BL.1RS	Quantum 7424	1997
1AL.1RS	Quantum 579	1994	1BL.1RS	Cougar	2000
1AL.1RS	TAM 110	1997	1BL.1RS	Wendy	2004
1AL.1RS	Hondo	1998	1BL.1RS	Endurance	2004
1AL.1RS	Nuhorizon	2000	1BL.1RS	Keota	2005
1AL.1RS	Above	2001	1BL.1RS	Shocker	2005
1AL.1RS	AP502 CL	2001	1BL.1RS	Art	2006
1AL.1RS	Charter	2004	1BL.1RS	Hitch	2008
1AL.1RS	Fannin	2004	1BL.1RS	Robidoux	2010
1AL.1RS	TAM 112	2005	1BL.1RS	SY Wolf	2010
1AL.1RS	TAM 303	2005	1BL.1RS	Gallagher	2012
1AL.1RS	Aspen	2006	1BL.1RS	Ruth	2015
1AL.1RS	Darrell	2006	1BL.1RS	Larry	2016
1AL.1RS	T140	2007			
1AL.1RS	Armour	2008			
1AL.1RS	T151	2008			
1AL.1RS	T153	2008			
1AL.1RS	T154	2008			
1AL.1RS	T158	2009			
1AL.1RS	Judee	2011			
1AL.1RS	TAM 305	2012			
1AL.1RS	AG Robust	2014			

beneficial effects to recipient wheats, including greater biomass (Shearman et al. 2005; Foulkes et al. 2007), increased heterosis in hybrid wheats (Owuoche et al. 2003), and increased root size (Waines and Ehdaie 2007). Thus, 1RS may convey to newly developed breeding lines enough of a yield advantage, in at least some backgrounds and environments, that breeders are unconsciously selecting and maintaining it. 1RS also might be maintained in breeding programs

because it *is* common in those elite lines that may be selected as parents. Inspection of pedigrees of lines entered in the SRPN and NRPN (USDA 2018) indicates most are derived from matings of experimental breeding lines, before any such lines are released as cultivars. Breeders select the best lines from their own and their colleagues' advanced trials, and enter them into their crossing blocks, without waiting to see if said lines are to be released or not. The

idea is to pyramid useful genes, and, in doing so, 1RS likely is present, perpetuating it even if the lines carrying it fail to achieve cultivar status.

Lukaszewski (2015) described 1BL.1RS as the “poster child” of alien introgressions. In the Great Plains of North America, it is joined and even exceeded by 1AL.1RS. The success of both translocations in North American winter wheat cultivar development, as described herein, and in both winter and spring wheats world-wide (Crespo-Herrera et al. 2017) highlight the value of alien introgressions, as proposed by McFadden and Sears (1947) many decades ago.

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#### Compliance with ethical standards

**Conflict of interest** The authors declare no conflict of interest.

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