

Sleeping Beauty Transposon Mutagenesis as a Tool for Gene Discovery in the NOD Mouse Model of Type 1 Diabetes

Colleen M. Elso,^{*,1} Edward P. F. Chu,^{*,†,1} May A. Alsayb,^{*,†,1} Leanne Mackin,^{*} Sean T. Ivory,^{*} Michelle P. Ashton,^{*,†,2} Stefan Bröer,[‡] Pablo A. Silveira,[§] and Thomas C. Brodnicki^{*,†,3}

^{*}Immunology and Diabetes Unit, St. Vincent's Institute, Fitzroy, Victoria 3065, Australia, [†]Department of Medicine, University of Melbourne, Parkville, Victoria 3010, Australia, [‡]Research School of Biology, The Australian National University, Canberra, ACT 2601, Australia, and [§]Dendritic Cell Research, ANZAC Research Institute, Concord, New South Wales 2139, Australia

ABSTRACT A number of different strategies have been used to identify genes for which genetic variation contributes to type 1 diabetes (T1D) pathogenesis. Genetic studies in humans have identified >40 loci that affect the risk for developing T1D, but the underlying causative alleles are often difficult to pinpoint or have subtle biological effects. A complementary strategy to identifying “natural” alleles in the human population is to engineer “artificial” alleles within inbred mouse strains and determine their effect on T1D incidence. We describe the use of the *Sleeping Beauty* (SB) transposon mutagenesis system in the nonobese diabetic (NOD) mouse strain, which harbors a genetic background predisposed to developing T1D. Mutagenesis in this system is random, but a green fluorescent protein (GFP)-polyA gene trap within the SB transposon enables early detection of mice harboring transposon-disrupted genes. The SB transposon also acts as a molecular tag to, without additional breeding, efficiently identify mutated genes and prioritize mutant mice for further characterization. We show here that the SB transposon is functional in NOD mice and can produce a null allele in a novel candidate gene that increases diabetes incidence. We propose that SB transposon mutagenesis could be used as a complementary strategy to traditional methods to help identify genes that, when disrupted, affect T1D pathogenesis.

KEYWORDS

forward genetics
reverse genetics
Slc16a10
Serinc1
amino acid
transporter

Type 1 diabetes (T1D) is an autoimmune disease in which lymphocytes mediate the specific destruction of insulin-producing pancreatic β cells (Bluestone *et al.* 2010). Genetic studies in human populations have detected >40 genomic intervals that harbor T1D-associated alleles (Polychronakos and Li 2011; Pociot *et al.* 2010). However, identification of the underlying genes for these T1D loci and the biological effects of putative causative alleles is often difficult due to genetic heterogeneity and limited tissue availability (Polychronakos and Li 2011; Pociot *et al.* 2010). Instead, it

has proven useful to complement human genetic studies with strategies that not only aim to discover “naturally” occurring alleles but also engineer “artificial” null alleles in putative and novel candidate genes to determine their effect on disease pathogenesis in inbred animal models (Ermann and Glimcher 2012).

The nonobese diabetic (NOD) mouse strain, in particular, has been widely used to investigate T1D pathogenesis (Driver *et al.* 2011; Jayasimhan *et al.* 2014). NOD mice spontaneously develop T1D, and genetic studies have identified >40 murine T1D susceptibility loci [termed insulin-dependent diabetes (*Idd*) loci], several of which overlap human T1D susceptibility loci (Burren *et al.* 2011). Although congenic NOD mouse strains have confirmed the majority of these *Idd* loci, relatively few of the underlying genes and their causative alleles have been definitively identified (Araki *et al.* 2009; Hamilton-Williams *et al.* 2001; Hung *et al.* 2006; Kissler *et al.* 2006; Laloraya *et al.* 2006; McGuire *et al.* 2009; Razavi *et al.* 2006; Tan *et al.* 2010; Yamanouchi *et al.* 2007). It has become apparent, however, that the NOD mouse strain has a combination of rare alleles (e.g., *H2-Ab^{g7}*) and common alleles (e.g., *H2-E^{null}* and *B2m^a*) for different genes that interact and increase the risk for T1D (Driver *et al.* 2011; Ridgway *et al.* 2008). Intriguingly, some nondiabetic mouse strains harbor a more

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¹These authors contributed equally to the work.

²Present address: DFG Center for Regenerative Therapies, Technische Universität Dresden, Dresden 01307, Germany.

³Corresponding author: Thomas C. Brodnicki, St. Vincent's Institute, 41 Victoria Parade, Fitzroy, Victoria 3065, Australia. E-mail: tbrodnicki@svi.edu.au

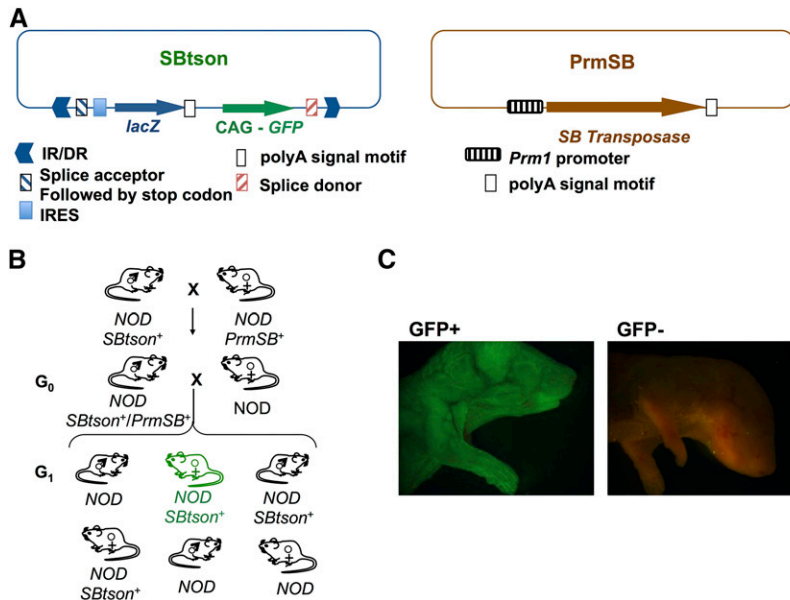


Figure 1 *Sleeping Beauty* transposon mutagenesis strategy. (A) Constructs used for production of NOD-*PrmSB* and NOD-*SBtson* lines. The transposon construct, pTrans-SA-IRESLacZ-CAG-GFP_SD:Neo, has been described (Horie *et al.* 2003). The transposase construct pRP1345 comprising *SB10* transposase driven by the mouse proximal protamine 1 promoter has also been described (Fischer *et al.* 2001). IR/DR: inverse repeat/direct repeat transposase recognition motifs. (B) Breeding scheme for *SB* transposon mutagenesis. NOD-*SBtson* mice (lines 1 and 2) were mated to NOD-*PrmSB* mice (lines 1 and 2). Double positive male offspring (seed mice) were backcrossed to wild-type NOD females to produce G₁ mice carrying potential transposon insertions. (C) Mice carrying transposon insertions that activated the polyA trap were detected by fluorescence under UV light prior to weaning.

diabetogenic allele than NOD mice for a given *Idd* locus (Brodnicki *et al.* 2003; Wang *et al.* 2014; Ghosh *et al.* 1993; McAleer *et al.* 1995). This complex genetic architecture for T1D susceptibility in the *Mus* species is similar to that described in humans and further complicates the identification of “natural” causative alleles within genes underlying *Idd* loci using traditional outcross and congenic mouse studies (Driver *et al.* 2011; Ridgway *et al.* 2008).

Here, we propose an alternative approach for disease gene discovery using the *Sleeping Beauty* (*SB*) transposon mutagenesis system to generate “artificial” alleles on the NOD genetic background. The *SB* transposon can insert within genes and disrupt transcript expression (Horie *et al.* 2003; Carlson *et al.* 2003; Ivics *et al.* 1997). Its unique sequence also serves as a molecular tag to rapidly identify the site of insertion without additional breeding. Transposition (*i.e.*, transposon “jumping”) is catalyzed by the *SB* transposase, which can be expressed *trans* and controlled by tissue-specific promoters to restrict transposition and subsequent gene mutation to germline or somatic cells. Once activated, transposition is relatively random, requiring only a target TA dinucleotide integration site and exhibiting some bias toward “jumping” in *cis*, *i.e.*, within the same chromosome (Keng *et al.* 2005; Carlson *et al.* 2003; Horie *et al.* 2003). A major advantage of the *SB* transposon is its ability to carry gene-trap elements and reporter genes, which increase gene disruption efficiency and accelerate identification of mice with a disrupted gene (Izsvak and Ivics 2004; Horie *et al.* 2003). Due to these unique characteristics, this system has been successfully used to mutate and characterize both putative and novel genes in different mouse models of cancer (Moriarity and Largaespada 2015; Dupuy *et al.* 2009). We show here that a relatively small-scale mutagenesis screen using the *SB* transposon, combined with disease-specific prioritization criteria, is able to identify a novel candidate gene that contributes to T1D susceptibility in NOD mice.

MATERIALS AND METHODS

Constructs and production of transgenic and *SB* transposon mutant mice

The transposase construct pRP1345 (Figure 1A) was obtained from Prof. R. Plasterk (Hubrecht Laboratory, The Netherlands) and has been previously described (Fischer *et al.* 2001). The transposon construct,

pTrans-SA-IRESLacZ-CAG-GFP_SD:Neo (Figure 1A), was obtained from Prof. J. Takeda (Osaka University, Japan) and has been previously described (Horie *et al.* 2003). *SB* transposon and *SB* transposase transgenic mice were produced by pronuclear injection of linearized constructs into fertilized NOD/Lt (NOD) oocytes at The Walter and Eliza Hall Institute Central Microinjection Service using a previously described protocol (Marshall *et al.* 2004). Transposon transgenic mouse lines are NOD-*TgTn(sb-Trans-SA-IRESLacZ-CAG-GFP-SD:Neo)1Tcb* and NOD-*TgTn(sb-Trans-SA-IRESLacZ-CAG-GFP-SD:Neo)2Tcb*, but have been termed NOD-*SBtson* L1 and L2 in the text. Transposase transgenic mouse lines are NOD-*Tg(Prm-sb10)1Tcb* and NOD-*Tg(Prm-sb10)2Tcb*, but have been termed NOD-*PrmSB* L1 and L2 in the text. NOD-*SBtson* mice were mated to NOD-*PrmSB* mice and double-positive hemizygous male offspring (NOD-*SBtson*⁺/*PrmSB*⁺) were identified by PCR genotyping. NOD-*SBtson*⁺/*PrmSB*⁺ males were backcrossed to wild-type NOD females to produce G₁ mice, carrying potential transposon insertions. Mice carrying transposon insertions, which activated the polyA trap, were noninvasively identified by visualizing GFP expression in newborn mice under UV light with confirmation by the presence of GFP fluorescence in ear biopsies using a fluorescent stereomicroscope equipped with a UV filter. Experiments involving mice were approved by the St. Vincent’s Institute Animal Ethics Committee.

Transposon copy number analysis

Southern blots were performed using standard protocols. An 898-bp probe specific for GFP labeled with α-³²P-dCTP using the DECA prime II Random Priming DNA labeling kit (Ambion) was used to detect the presence of the *SB* transposon. Copy number was calculated by comparison with standards of known copy number.

Transposon insertion site identification

Ligation-mediated PCR (LM-PCR) was performed based on previously described protocols (Largaespada and Collier 2008; Takeda *et al.* 2008; Horie *et al.* 2003; Devon *et al.* 1995). Briefly, 1 μg genomic DNA was digested with *Hae*III, *Alu*I, *Bfa*I, or *Nla*III (New England Biolabs), and splinkerettes (Table 1) compatible with appropriate blunt or cohesive ends were ligated to the digested genomic DNA fragments using T4 DNA ligase (New England Biolabs). The two oligonucleotides to

■ **Table 1** Oligonucleotides used as splinkerettes and primers for LM-PCR

Name ^a	Sequence	Purpose ^a
SplB-BLT	CGAATCGTAACCGTTTCGTACGAGAATCGCTGTCTCTCCAACGAGCCAAGG	Splinkerette
Spl-top	CCTTGGCTCGTTTTTTTGCAGAAA	Splinkerette
Nla linker+	GTAATACGACTCACTATAGGGCTCCGCTTAAGGGACCATG	Splinkerette
Nla linker-	5'-GTCCCTTAAGCGGAGCC-amino	Splinkerette
Bfa linker+	GTAATACGACTCACTATAGGGCTCCGCTTAAGGGAC	Splinkerette
Bfa linker-	5'-TAGTCCCTTAAGCGGAG-amino	Splinkerette
Spl-P1	CGAATCGTAACCGTTTCGTACGAGAA	LM-PCR
Spl-P2	TCGTACGAGAATCGCTGTCTCTCC	Nested LM-PCR
T/JBA	TAAGTACCTTAAGACAGGGAATCTTTAC	LM-PCR (left)
TJB1	TTTACTCGGATTAATGTCAGGAATG	Nested LM-PCR (left)
TDR2	CTGGAATTGTGATACAGTGAATTATAAGTG	LM-PCR (right)
T/BAL	CTTGTGTCATGCACAAAGTAGATGTC	Nested LM-PCR (right)
LongIRDR(L2)	CTGGAATTTTCCAAGCTGTTTAAAGGCACAGTCAAC	LM-PCR (left)
NewL1	GACTTGTGTCATGCACAAAGTAGATGTC	Nested LM-PCR (left)
LongIRDR(R)	GCTTGTGGAAGGCTACTCGAAATGTTTGACCC	LM-PCR (right)
KJC1	CCACTGGGAATGTGATGAAAGAAATAAAGC	Nested LM-PCR (right)
Linker primer	GTAATACGACTCACTATAGGGC	LM-PCR
Linker primer nested	AGGGCTCCGCTTAAGGGAC	Nested LM-PCR

^a Primer names and purpose are based on the previously described LM-PCR protocols (Keng et al. 2005; Largaespa and Collier 2008).

produce the double-stranded splinkerette were annealed at a concentration of 50 μ M in the presence of 100 mM NaCl by incubating at 95 $^{\circ}$ for 5 min and then allowing the mixture to cool to room temperature in a heat block before ligation. A second digest, with *Xho*I or *Kpn*I, was used to remove ligated splinkerettes from transposon concatemer fragments. DNA was purified after each step using QiaQuick PCR Purification kit (Qiagen). Two rounds of PCR were performed using nested primers within the linker and transposon (Table 1) with the subsequent PCR product sequenced. Restriction enzymes, splinkerettes, and primers were used in the following six combinations (first digest; splinkerette; second digest; primer set 1; primer set 2): (i) and (ii) *Alu*I or *Hae*III; SplB-BLT; *Xho*I; T/JBA \times Spl-P1; TJB1 \times Spl-P2; (iii) and (iv) *Alu*I or *Hae*III; SplB-BLT; *Kpn*I; TDR2 \times Spl-P1; T/BAL \times Spl-P2; (v) *Bfa*I; Bfa linker; *Kpn*I; LongIRDR(L2) \times Linker primer; NewL1 \times linker primer nested; and (vi) *Nla*III; Nla linker; *Xho*I; LongIRDR(R) \times Linker primer; KJC1 \times Linker primer nested. The resulting sequence was aligned to mouse genome build GRCm38 to identify the transposon-flanking genomic sequence (*i.e.*, insertion site) and determine which gene was disrupted based on current annotation for build GRCm38. The different combinations give a number of chances to identify genomic DNA adjacent to both the 3' and 5' ends of the transposon after transposition. Additional primers were designed for genotyping the transposon insertion site in established mutant mouse lines (Table 2).

Gene expression analysis

RNA was extracted from tissues using Trizol (Invitrogen) and cDNA was synthesized using Superscript III (Invitrogen), both according to manufacturer's instructions. For SB4 mice, RT-PCR was performed with oligos specific for *Slc16a10*: Exon 1 forward: 5'-GTGGTGCAACGGGTCGGTGT; Exon 2 reverse: 5'-GACACTCACGATGGGCAGCA; Exon 6 reverse: 5'-ACAGAGCACAACCCCCAACG; and *Actb* (forward: 5'-CGGTTCCGATGCCCTGAG; reverse: 5'-TGATCACATCTGCTGGAAGG). For SB7 mice, RT-PCR was performed with primers in *GFP* (5'-CCCTGAGCAAAGACCCCAACGAGAAGC) and *Serin1* exon 1 (reverse: 5'-TCGTCCTTTTCTGGGCTTA) or exon 2 (reverse: 5'-ACTCCGACGAGCAAGAAAAG), followed by Sanger sequencing of products. Transposase expression was determined using quantitative real-time PCR using LightCycler Probe Master Reagent

(Roche Diagnostics) and the following primer/UPL probe combination for SB transposase (*SB10*): Fwd: 5'-ACCACGCAGCCGTCATAC; Rev: 5'-CACCAAAGTACGTTTCATCTCTAGG; UPL probe #12: GGAAGGAG; and *Hprt*: Fwd: 5'-TCCTCCTCAGACCGCTTTT; Rev: 5'-CCTGGTTCATCATCGCTAATC; UPL probe #95: AGTCCCAG. Data were normalized to *Hprt* expression. For SB7, *Serin1* expression was determined by quantitative real-time PCR (LightCycler480, Roche) with the following primer/probe combination for *Serin1*: (Fwd: 5'-GACGCGCGGCGGAT; Rev: 5'-GCATCGGCACAGCAAACAC; Taqman Probe: 5'-GCTGGATTCGGTGT) and normalization was performed using *Hprt* (Taqman assay: Mm01545399_m1). Data were normalized to *Hprt* expression.

Diabetes monitoring

Mice were tested once per week for elevated urinary glucose using Diastix reagent strips (Bayer Diagnostics). Mice with a positive glycosuria reading (>110 mmol/L) and confirmed by a positive glucose reading (>15 mmol/L), using Advantage II Glucose Strips (Roche), were diagnosed as diabetic.

Data availability

Mouse lines are available upon request.

RESULTS

Generation of transposition events in NOD mice using SB transgenic NOD lines

To perform germline mutagenesis of the NOD mouse, we used two SB constructs previously used in mice (Figure 1A). The pRP1345 construct contains the transposase gene under the control of the proximal protamine 1 (*Prm1*) promoter, which restricts expression to spermatogenesis and limits transposon mutations to the germline, thus preventing somatic mutations (Fischer et al. 2001). The SB transposon construct (*sb-pTrans-SA-IRESLacZ-CAG-GFP-SD:Neo*) contains splice acceptor and donor sequence motifs encompassing a promoter trap comprising the *lacZ* gene, and a polyA trap with the gene encoding enhanced green fluorescent protein (*GFP*) driven by the CAG promoter (Horie et al. 2003). This construct enables efficient identification of mutant mice in

■ **Table 2** Oligonucleotides used as primers for genotyping SB4 and SB7 mice

Line	Fwd	Rev	Size	Allele
SB4	AGCCCAGAAGACAACCCCTCTTGT	AAAGGGGGCGTGCGCTAAACA	123 bp	WT
SB4	CTTGTGTCATGCACAAAGTAGATGTCC	AAAGGGGGCGTGCGCTAAACA	224 bp	SB
SB7	ATTCCACCAGTGATGTGCTGGTAAC	CCTTGAAATCATCCCGTGAGAGA	647 bp	WT
SB7	ATTCCACCAGTGATGTGCTGGTAAC	CTTGTGTCATGCACAAAGTAGATGTCC	632 bp	SB

which the transposon has inserted in a gene, as activation of the polyA trap results in ubiquitous GFP expression, *i.e.*, mutant mice fluoresce.

The transposase and transposon were introduced separately into NOD mice to establish independent transgenic NOD lines for each SB component. Briefly, SB transposon and SB transposase transgenic mice were produced by pronuclear injection of linearized constructs into fertilized NOD/Lt (NOD) oocytes. NOD-*Tg(Prm-sb10)* mice (called NOD-*PrmSB* hereafter) harbor the transposase; and NOD-*TgTn(sb-pTrans-SA-IRESLacZ-CAG-GFP-SD:Neo)* mice (called NOD-*SBtson* hereafter) harbor the transposon. The breeding scheme to generate mutant NOD mice is outlined in Figure 1B: mice from the two transgenic lines are mated, bringing together the two components of the SB system, and transposition occurs within the sperm cells of the double-positive hemizygous “seed” males. These NOD seed males are mated to wild-type NOD females to generate G₁ litters. G₁ pups with potential transposon-disrupted genes are efficiently identified before weaning by visualization of bodily GFP expression under UV light (Figure 1C). This early detection allows cage space to be minimized; only those litters that contain fluorescent pups are weaned and kept for additional analysis.

To assess the feasibility of the SB mutagenesis system in the NOD mouse strain, two NOD-*SBtson* lines were established for which the number of copies of the transposon within the transgene concatemer was determined by Southern Blot analysis (Figure 2A) and the site of transgene integration was determined by ligation-mediated PCR (LM-PCR) (Figure 2B). These lines were bred with two established NOD-*PrmSB* lines, in which expression of SB transposase in testes had been confirmed by quantitative RT-PCR (data not shown), to produce seed males. NOD seed males (*i.e.*, double-positive, hemizygous for both SB constructs) were backcrossed to NOD females to produce G₁ mice. Three different breeding combinations of transposon/transposase transgenic strains gave rise to GFP-positive offspring at the rate of 2.0%, 2.3%, and 2.7% respectively (Figure 2B). A fourth combination did not produce any mutant pups (0%), but breeding of this fourth combination was stopped before reaching a similar total of G₁ mice to the other combinations (Figure 2B). Eleven GFP-positive G₁ mice were sired, confirming that these transgenic lines can facilitate SB transposition events on the NOD genetic background and resultant GFP-positive mice can be detected.

Identification and prioritization of transposition sites in GFP-positive G₁ NOD mice

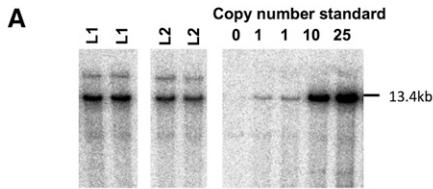
LM-PCR followed by sequencing and genome alignment was used to identify the transposition sites for nine of the 11 GFP-positive mice (Table 3). Of the two GFP-positive mice that were not determined, one died before characterization. It was not clear why the second GFP-positive mouse was refractory to site identification by LM-PCR despite using six different restriction digest/PCR combinations. Consistent with the published >30% rate of local chromosomal hopping (Keng *et al.* 2005), seven of the nine identified transposition sites fell on the same chromosome as the transposon donor concatemer. Of the seven insertion sites derived from the NOD-*SBtsonL1* concatemer on chromosome (Chr) 10, five were on Chr10, with the other two identified on Chr1 and Chr4. The two insertion sites arising from the

NOD-*SBtsonL2* concatemer on Chr1 were both mapped to Chr1. LM-PCR also indicated that each GFP-positive mouse contained a single, rather than multiple, transposon insertion site.

Once GFP-positive NOD mice and their transposon mutations are identified, there are two options. One option is to generate and monitor diabetes onset in cohorts of mice from every GFP-positive G₁ mouse. Although such a full-scale phenotype-driven approach is aimed at identifying novel genes not suspected to play a role in diabetes, as well as known and putative candidate genes, this option requires substantial animal housing capacity and monitoring numerous cohorts for diabetes over a >200-d time course. Like many investigators, we have limited resources and this option was not feasible. However, the SB transposon mutagenesis strategy allows for a second option: prioritizing mutant mice based on the genes that are disrupted and establishing homozygous mutant lines for expression and diabetes monitoring. We therefore prioritized transposon-disrupted genes based on the following criteria:

- 1) The transposon insertion site is predicted to disrupt the gene in some way. Has the transposon inserted within an exon, an intron, or in a regulatory region? Is the mutation predicted to disrupt gene expression? An insertion that is predicted to completely abrogate expression of the normal gene product would be of high priority; however, in some instances a predicted hypomorphic allele may also be of value.
- 2) The gene is a known or putative candidate for a described mouse and/or human T1D susceptibility locus. T1D loci and candidate genes are curated in a searchable form at T1Dbase (<https://t1dbase.org/>) (Burren *et al.* 2011).
- 3) The gene, not previously considered as a putative T1D susceptibility gene, encodes a known protein involved in an immune-related molecular pathway that could affect T1D pathogenesis (*e.g.*, cytokine or chemokine signaling/regulation, pattern recognition pathways, costimulatory molecules, apoptosis, regulation of immune tolerance), but is not likely required for general immune cell development.
- 4) The gene, not previously considered as a putative T1D susceptibility gene, is expressed in relevant cells (*e.g.*, immune cell subsets, β cells) as determined in the first instance using gene expression databases [*e.g.*, Immunological Genome Project (<https://www.immgen.org/>)] (Heng *et al.* 2008) followed by additional expression analyses if needed.

Of the nine transposon insertion sites identified (Table 3), only four were located within or nearby (within 5 kb) annotated genes or expressed sequence tags (ESTs): SB4 in intron 1 of *Slc16a10*; SB7, 4.6 kb 5' of *Serinc1*; SB8, 5 kb 5' of *AKO18981*; and SB9 in intron 2 of *Pard3b*. In particular, the transposon within *Pard3b* inserted in the opposite orientation to the direction of transcription of the gene, so it seems unlikely that it would disrupt expression of *Pard3b*, which encodes a cell polarity protein most highly expressed in the kidney (Kohjima *et al.* 2002). There is, however, an antisense gene, *Pard3bos1*, that overlaps *Pard3b* approximately 102 kb downstream of the transposon insertion site. Thus, it is more likely that the polyA gene-trap has



B Characterization of NOD-SBtson lines

Transgenic line	Copy ^a	Site of Transgene Integration chr:bp	x NOD- <i>PrmSBL1</i> ^b		x NOD- <i>PrmSBL2</i> ^c	
			# pups	GFP ⁺ (%)	# pups	GFP ⁺ (%)
NOD-SBtsonL1	5	chr10:54,651,092	176	4 (2.3)	252	5 (2.0)
NOD-SBtsonL2	3	chr1:49,283,271	58	0 (0.0)	75	2 (2.7)

Figure 2 Characterization of NOD-SBtson lines and frequency of GFP-positive G₁ pups. (A) Transgene copy number in NOD-SBtson lines was determined by Southern blot analysis using standard techniques and an 898-bp probe specific for *GFP* that detected a 13.4-kb band containing the transposon. Known amounts of transposon DNA were used to generate a standard curve for determining copy number. (B) Site of transgene integration was determined by LM-PCR (coordinates are based on genome build GRCm38). Number and percentage of GFP⁺ offspring for each breeding combination is shown. ^aNumber of copies of the transposon in the donor concatemer, determined by Southern blot as shown in (A). ^bOffspring generated from seed males produced by breeding NOD-SBtson lines with NOD-*PrmSBL1*. ^cOffspring generated from seed males produced by breeding NOD-SBtson line with NOD-*PrmSBL2*.

been activated by splicing into this antisense gene. The remaining five transposons inserted in regions currently without any annotated genes, suggesting either the presence of unannotated genes or the presence of cryptic polyA sequence motifs as the GFP polyA trap was activated.

The following observations were made regarding the prioritization of the nine identified transposon insertion sites. Only one fell within a T1D susceptibility locus (*Pard3bos1* in *Idd5.1*) (Burren *et al.* 2011), but unfortunately this mouse (SB9) died before it could be bred. *AK018981* has no publicly available information except that it was sequenced from RNA isolated from adult mouse testes. Both *Serinc1* and *Slc16a10* have been previously disrupted in mice: *Serinc1* on a mixed C57BL/6 × 129 genetic background (Tang *et al.* 2010) and *Slc16a10* on a C57BL/6 background (Mariotta *et al.* 2012). Neither knockout mouse strain

was reported to develop spontaneous disease. The strains either have not been analyzed for immunological phenotypes (*Slc16a10*) or have only been analyzed in small cohorts ($n \leq 4$) with no differences observed (*Serinc1*). Despite this, *Serinc1* and *Slc16a10*, although not directly attributed in the literature with immune-related roles, have functions that could be postulated to affect immune cell responses and/or β -cell activity, which could be revealed in the context of the “sensitized” NOD genetic background (*i.e.*, the NOD mouse strain enables detection of mutated genes that increase or decrease diabetes incidence). Notably, both genes according to expression databases (The Immunological Genome Project and BioGPS) (Heng *et al.* 2008; Wu *et al.* 2009) were highly expressed in macrophages, an immune cell population with a key role in T1D pathogenesis (Driver *et al.* 2011; Jayasimhan *et al.* 2014). SB4 and SB7, which were both male G₁ mice, did not carry the transposase construct; consequently, secondary jumping of the transposon was not possible in their offspring. SB4 and SB7 G₁ mice were thus prioritized for establishment of new lines, bred to homozygosity, and investigated for the effect of their transposon insertions.

Characterization of transposon effects in two prioritized GFP-positive G₁ NOD mice

The transposon insertion site in the SB7 mouse (NOD. *Serinc1*^{Tn(sb-Trans-SA-IRESLacZ-CAG-GFP-SD;Neo)1.7Tcb}) localized 4.6 kb upstream of *Serinc1* (Table 3, Figure 3A). SERINC1 facilitates the synthesis of serine-derived lipids, including the essential membrane lipids phosphatidylserine and sphingolipid (Inuzuka *et al.* 2005). These are important components of membrane structures known as “ordered membrane domains” or “lipid rafts,” which are required for appropriate signaling in immune cells (Szoor *et al.* 2010; Yabas *et al.* 2011). Due to the position of the transposon insertion, we postulated that, rather than completely disrupting expression of *Serinc1*, the mutation may affect gene regulation. RT-PCR analysis of homozygous SB7 splenic RNA using primers within *GFP* and *Serinc1* identified two fusion transcripts in addition to the normal transcript. The first contains *GFP* spliced to sequence upstream of exon 1 with normal splicing of the entire gene. The second contains *GFP* spliced directly to exon 2 of *Serinc1* (Figure 3A). The skipped exon 1 encodes the first 13 amino acids of the *Serinc1* coding sequence. The production of any SERINC1 protein from these fusion transcripts is unlikely because there is a stop codon following the *GFP* sequence and no obvious internal ribosomal entry site prior to the *Serinc1* sequence, but this still remains to be tested. In either case, the presence of the transposon

Table 3 Sleeping Beauty transposon insertion sites in GFP-positive G₁ NOD mice

Mutant Mouse	NOD-SBtson Line	NOD- <i>PrmSB</i> Line	Site of Transposion Insertion (STI) ^a	Closest Gene in Correct Orientation	Gene Coordinates ^a	Position of STI with Respect to Gene
SB1	L1	L1	chr4:19,247,537	<i>Cnbd1</i>	chr4:18,860,454-19,122,526	125 kb 5' of gene
SB2	L1	L1	chr10:30,040,682	<i>Rspo3</i>	chr10:29,453,107-29,535,867	505 kb 5' of gene
SB3	L1	L1	chr10:54,845,255	<i>Msl3l2</i>	chr10:56,106,917-56,116,880	1.2 Mb 5' of gene
SB4	L1	L1	chr10:40,122,645	<i>Slc16a10</i>	chr10:40,033,535-40,142,254	Intron 1
SB5 ^c	L1	L2	n.d.	n.d.	n.d.	n.d.
SB6	L1	L2	n.d.	n.d.	n.d.	n.d.
SB7	L1	L2	chr10:57,537,126	<i>Serinc1</i>	chr10:57,515,775-57,532,529	4.6 kb 5' of gene
SB8 ^c	L1	L2	chr10:56,499,234	<i>AK018981</i>	chr10:56,504,501-56,505,287	5 kb 5' of gene
SB9 ^c	L1	L2	chr1:61,954,348	<i>Pard3bos1</i>	chr1:61,767,415-61,851,462	102 kb 5' of gene ^b
SB10	L2	L2	chr1:49,094,718	<i>C230029F24Rik</i>	chr1:49,244,616-49,340,431	150 kb 5'
SB11	L2	L2	chr1:48,800,150	<i>Slc39a10</i>	chr1:46,807,544-46,853,509	1.9 Mb 5' of gene

^a Coordinates are based on genome build GRCm38.

^b The SB9 transposon site of integration falls within intron 2 of *Pard3b* in the opposite orientation.

^c SB5, SB8, and SB9 mice died of unknown causes before homozygous lines could be established. n.d., not determined.

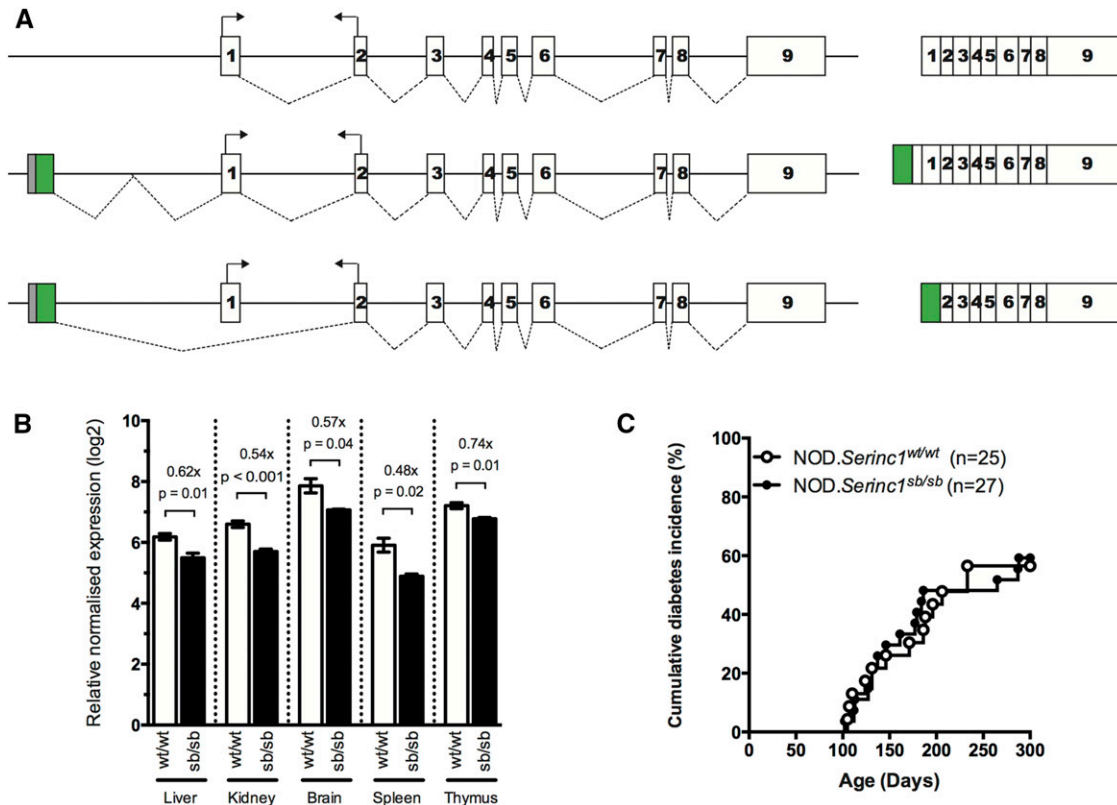


Figure 3 Analysis of *Serinc1* expression and diabetes incidence in SB7 mutant mice. (A) Schematic diagrams of *Serinc1* gene and transcripts with transposon insertion. *Serinc1* consists of nine exons and gives rise to a 2889-bp spliced transcript (top diagram). The transposon insertion occurs 4.6 kb 5' of *Serinc1*. RT-PCR led to the identification of two fusion transcripts: one that splices from GFP (green) to sequence upstream of exon 1 resulting in otherwise normal splicing of *Serinc1* (middle diagram), and another that skips exon 1 (bottom diagram). (B) Expression analysis was performed using RNA isolated from tissues of wild-type (wt/wt) and homozygous mutant (sb/sb) littermates (n = 4). Expression was determined by quantitative real-time PCR. Exon locations of *Serinc1* primers are indicated by arrows in (A). Error bars represent \pm SEM. Statistical significance for the difference in expression was obtained using pairwise t-tests. (C) The cumulative diabetes incidence was determined for age-matched female cohorts monitored concurrently. Pairwise comparisons of diabetes incidence curves were performed using the log-rank test.

insertion results in a relatively small, but significant, decrease in expression of *Serinc1* as measured by quantitative RT-PCR using a Taqman probe spanning the exon 1/2 splice junction (Figure 3B). This splice junction is used in both the “normal” *Serinc1* and the fusion transcript that includes exon 1; therefore, the reduction in “normal” *Serinc1* expression is greater than measured by this assay. These analyses suggest that, rather than completely disrupting expression of *Serinc1*, this mutation modulates expression and represents a hypomorphic allele. Nonetheless, homozygous mutant SB7 mice exhibited a similar diabetes incidence compared to wild-type littermate females (Figure 3C), indicating that a minor reduction in *Serinc1* expression does not affect T1D pathogenesis.

The transposon insertion identified in the SB4 mouse (NOD. *Slc16a10*^{Tn(sb-Trans-SA-IRESLacZ-CAG-GFP-SD:Neo)1.4Tcb}) localizes to intron 1 of *Slc16a10* (Table 3, Figure 4A), which encodes the aromatic amino acid transporter SLC16A10 (also known as TAT1) (Mariotta *et al.* 2012; Ramadan *et al.* 2006). It is becoming increasingly evident that regulation of amino acid transport is crucial for the proper regulation of immune cell activation and function (Nakaya *et al.* 2014; Sinclair *et al.* 2013; Thompson *et al.* 2008), and also impacts glycemic control (Jiang *et al.* 2015). We predicted that the strong splice acceptor encoded by the transposon would result in splicing from exon 1 of *Slc16a10* into the transposon sequence, thus truncating the *Slc16a10* transcript. Expression analysis of homozygous mutant SB4 mice showed that the

Slc16a10 transcript was not detected (Figure 4B), which was associated with a significant increase in diabetes incidence compared to wild-type littermate females (Figure 4C). This result demonstrates that SB transposon mutagenesis can be used to identify novel genes that affect T1D pathogenesis. Moreover, SB4 is a promising mutant mouse line for investigating the role of *Slc16a10* and aromatic amino acid transport in macrophage function and the development of T1D in NOD mice.

DISCUSSION

We demonstrate here that SB transposon mutagenesis can successfully generate both null and hypomorphic mutations in the NOD mouse. A significant advantage of this strategy is the use of the GFP reporter to screen out, before weaning, those mice unlikely to be carrying a functional mutation, thereby significantly reducing mouse handling and housing. Mutation sites can then be determined in individual GFP-positive mice and prioritized for further analysis based on the requirements of the investigator. We used our prioritization criteria to select two mutant NOD mice (SB4 and SB7) for further characterization and subsequently found that disruption of *Slc16a10* expression in NOD mice resulted in an increased T1D incidence. Although further studies are required to determine how *Slc16a10* and amino acid transport contributes to diabetes pathogenesis, this result indicates that SB transposon mutagenesis can be used as a complementary approach to other T1D gene discovery strategies.

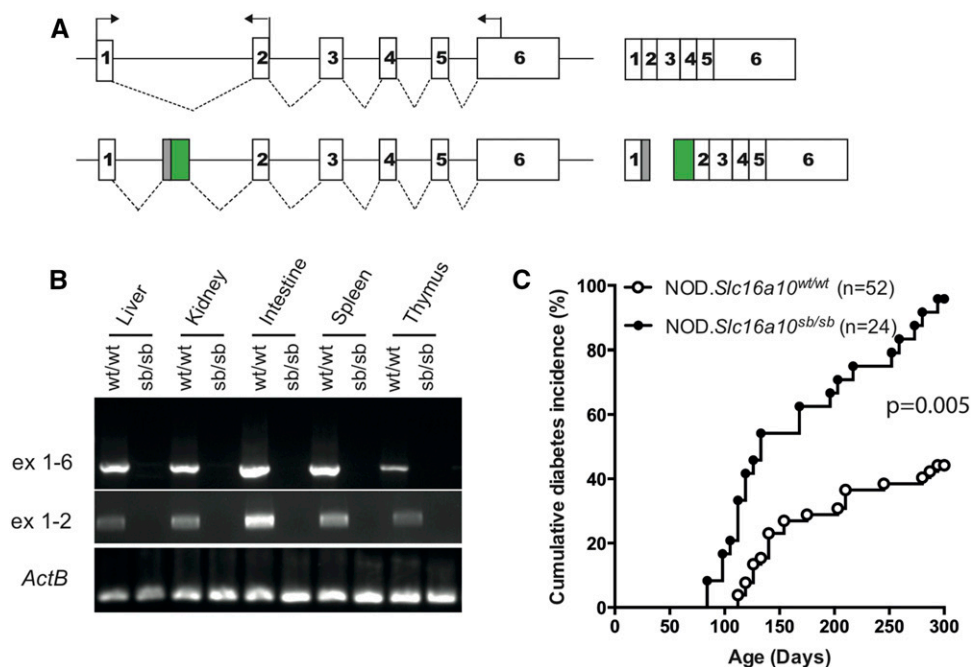


Figure 4 Analysis of *Slc16a10* expression and diabetes incidence in SB4 mutant mice. (A) Schematic diagrams of *Slc16a10* gene and transcript with transposon insertion. *Slc16a10* consists of six exons and gives rise to a 5394-bp spliced transcript (top diagram). The transposon insertion occurs within intron 1 and is predicted to prevent splicing between exons 1 and 2, instead generating a truncated transcript [exon 1 - transposon (gray box)] and a transcript initiated by the CAG promoter and GFP gene [transposon (green box) - exon 2] (bottom diagram). (B) Detection of transcript expression was performed by RT-PCR using RNA isolated from tissues of wild-type (wt/wt) and homozygous mutant (sb/sb) littermates. Spliced products for exon 1/exon 2 or exon 1/exon 6 were not detected in any tissues tested from homozygous mutant mice. Expression analysis is representative of three mice per genotype. (C) The cumulative diabetes incidence was determined for age-matched female cohorts monitored concurrently. Pairwise comparisons of diabetes incidence curves were performed using the log-rank test.

We observed 2–3% fluorescent G_1 offspring, which is lower than that reported for a similar mutagenesis scheme using the same transposon construct (~7%) (Horie *et al.* 2003). If one aims for a single transposition event per G_1 offspring, it would be expected that ~20% of the G_1 offspring should fluoresce [*i.e.*, ~40% of the mouse genome contains exons/introns (Sakharkar *et al.* 2005), with a 50% chance of the transposon landing in the correct orientation to trap the polyA sequence motif]. There are a number of possibilities that could explain a lower rate of fluorescent G_1 offspring, pointing to improvements that can be made to our screen. The NOD-*SBtsn* “mutator” transgenic lines we generated harbored few copies of the transposon. A higher copy number in the donor transposon concatemer may increase mutation efficiency due to more “jumping” transposons in the sperm of NOD seed males (Geurts *et al.* 2006). Use of a more efficient transposase could also increase transposition efficiency. As the first described transposon for use in vertebrates, the *SB* system has been the most widely used and developed. Improvements from the first-generation *SB* transposases, such as that used in our study, have seen 100-fold increases in transposition efficiency (Mates *et al.* 2009).

Trying to obtain too high of a mutation rate, however, is not necessarily favorable. For example, increasing transposon copy number by using transgenic mice with large transposon concatemers (>30 copies) may lead to local chromosomal rearrangements in subsequent offspring and complicate characterization of causative transposon mutations (Geurts *et al.* 2006). Increasing transposon number and/or transposase efficiency will also lead to GFP-positive offspring with multiple gene mutations. It would then require additional work to identify and confirm all the gene mutations in a given mouse, as well as additional breeding to segregate mutations and test mutant mice with only one gene mutation, all of which increases breeding times

and cage costs. Thus, an advantage of a lower efficiency is that it leads to mice with only one gene mutation, eliminating the need for extensive segregation analysis and facilitating more efficient gene identification and prioritization. Although we did not establish and characterize all of our mutant mouse lines due to limited resources, cryopreservation of sperm from mutant male mice could be used to allow archiving of mutants for future investigation. Empirically, it will be up to an individual laboratory to determine how many offspring they can efficiently screen and how they will prioritize mutant mice for subsequent analysis based on the mutation rate of their transgenic lines and available resources.

Interestingly, several of our transposon insertion sites mapped to regions at some distance from annotated genes. Although it is possible that the GFP could be activated by splicing into a cryptic polyA site, polyA gene trap strategies have been successfully used to identify novel unannotated genes (Zambrowicz *et al.* 1998). However, it may be difficult to predict *a priori* if an unannotated gene will be of interest, especially if it has little to no sequence homology with known genes. In this regard, the *SB* mutagenesis approach enables a phenotype-driven (*i.e.*, forward genetics) approach to test novel genes that might not otherwise be targeted using a candidate gene approach (*i.e.*, reverse genetics). *SB* mutagenesis may also benefit characterization of regions (*e.g.*, *Idd* loci) that are known to contain putative susceptibility genes, but for which the “natural” causative alleles have not yet been identified. Saturation mutagenesis could effectively be performed in these regions by generating a transgenic NOD mouse line containing a transposon concatemer near the region of interest and taking advantage of the propensity of *SB* transposons to reintegrate close to the donor transposon concatemer (Keng *et al.* 2005). Nonetheless, investigating unannotated genes is riskier (*i.e.*, the disrupted gene may not affect T1D) or more costly in the case of saturation mutagenesis (*i.e.*, more mutant

lines need to be generated and characterized). Hence, prioritizing genes based on function postulated to contribute to T1D pathogenesis may be more favorable to most labs.

Transposon mutagenesis is one of a range of techniques that can be used to identify gene variants that affect development of T1D. Conventionally, “artificial” null alleles for candidate genes have been generated in other strains and bred onto the NOD background by serial backcrossing. This approach, however, results in the null allele being encompassed by a “hitchhiking” congenic interval from the other strain, which may also affect T1D susceptibility (Simpfendorfer *et al.* 2015; Armstrong *et al.* 2006; Leiter 2002; Kanagawa *et al.* 2000). Although NOD ES cell lines are available (Hanna *et al.* 2009; Nichols *et al.* 2009; Ohta *et al.* 2009), there are relatively few reports of their use in targeting genes (Kamanaka *et al.* 2009; Morgan *et al.* 2013). Alternatively, random mutagenesis using *N*-ethyl-*N*-nitrosourea (ENU) does not require ES cells or prior knowledge about candidate genes. ENU mutagenesis, however, creates tens to hundreds of mutations per mouse. Substantial breeding and sequencing, more than for SB transposon mutagenesis, would be required to segregate and test individual ENU mutations for their effect on T1D susceptibility (Hoyne and Goodnow 2006). Finally, emerging gene-editing techniques using the CRISPR-Cas9 system (Ran *et al.* 2013) can facilitate hypothesis-driven investigation of known and putative candidate genes in NOD mice (Ran *et al.* 2013; Li *et al.* 2014). Modifications of the CRISPR-Cas9 system also allow a range of strategies to be used, including the production of conditional alleles, insertion of reporters, activation or repression of alleles, and specific gene editing allowing the recreation of particular variants affecting immune function (Pelletier *et al.* 2015). Nonetheless, this approach requires genes to be specifically targeted, whereas SB transposon mutagenesis is random and may identify genes not otherwise considered. Thus, a combination of different approaches for gene discovery and characterization of allelic effects is available and will likely prove useful for understanding the genetic architecture of T1D.

The increasing number of causative “natural” alleles identified in human populations and inbred mouse strains will undoubtedly aid our understanding and prediction of genetic risk for T1D, as well as aid future clinical trials in selecting appropriate patient treatment groups based on their genetic profile (Bluestone *et al.* 2010; Polychronakos and Li 2011). Nonetheless, many of the identified T1D loci and underlying causative alleles have subtle biological effects that are not therapeutically amenable or are difficult to investigate due to tissue availability. Generating random “artificial” null alleles in the NOD mouse provides an alternative strategy to test and identify both putative and novel genes that: (i) have larger diabetes effects when more grossly disrupted; (ii) represent potential drug targets; and (iii) are less likely to be identified in population-based studies of natural variation. The NOD mouse exhibits a number of immunological abnormalities that are associated with T1D pathogenesis and provides a “sensitized” background to investigate the effect of artificial mutations upon the development of T1D (Driver *et al.* 2011; Ridgway *et al.* 2008; Jayasimhan *et al.* 2014). Our study indicates that SB transposon mutagenesis in NOD mice is feasible and provides a new strategy that combines the advantage of both forward genetics (random mutagenesis) and reverse genetics (gene prioritization) for the potential discovery of new genes that affect T1D pathogenesis.

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