

Genetic monitoring of breeding colony through evaluation of genetic diversity within and between the inbred lines by using microsatellite markers

Satheesh Kumar P and Govind Yadav

Animal House Division, CSIR-Indian Institute of Integrative Medicine, Jammu

Corresponding author:

Satheesh Kumar P

Animal House Division, CSIR-Indian Institute of Integrative Medicine,
Jammu-180001

Email : satheesh@iiim.ac.in

Abstract

BALB/c strain is the most commonly used animal research mouse model in biology/Bio-medical research. From BALB/c strain many of inbred, congenic and transgenic models were established and being used for the betterment of humans as well as domestic animals. At CSIR-IIIM, Jammu animal house facility conducted study for evaluation genetic diversity of inbred lines of BALB/c strain. Animals were maintained in SPF condition with routine health monitoring. Selection was conducted based on growth and reproduction performance of the mothers. Four selected lines and an unselected control line were maintained throughout the experiment. After, 20th generation of full sib mating, we obtained three different lines. All the three lines of BALB/c strain shows standardized litter size of 8 pups at birth. All line mice were genotyped by using established 14 pair of microsatellite markers and genetic diversity analysis. The Effective number of alleles was reported 1.021 ± 0.016 in overall experimental population. Shannon's Index for within lines was 0 and among lines was 0.019 ± 0.014 . The Estimated Diversity (u) was 0 for within lines, 0.052 ± 0.036 for control lines and 0.013 ± 0.009 for among population. Polymorphic information content (PIC) observed in D11Mit260 marker. Line1 was showing very little Genetic distance with Line 2 (0.154) followed by control line (0.114) and Line 3 (0.074). Control line and Line 2 showed very close genetic distance (0.010). It is concluded that BALB/c strain has no diversity within line and has very little diversity between lines after 20th generation of full-sib mating.

Keywords: Mouse, Litter Size, Genetic Diversity, Microsatellite Marker.

Introduction

Albino mice of a, b, c Stocks was acquired by Halsey Bag in 1913. In 1932, Snell added the 'c' to denominate albino (www.jax.org/). Now BALB/c mouse strain is widely distributed and amongst the top 2-3 most widely used inbred strains which is used as a general-purpose strain in many different disciplines. It has good breeding performance and long reproductive life-span. BALB/c strain is maintained by inbreeding (full sib or sister x brother mating) at CSIR-IIIM, Jammu. Four (or three?) selected lines and an unselected control line were maintained throughout the experiment. After 20th generation of full sib mating, we obtained three different lines. All the three lines of BALB/c strain shows standardized litter size of 8 pups at birth. Which are genotypically identical and have high level of homozygosity. For quality control CSIR-IIIM, Jammu has established numerous production procedures and

Genetic Monitoring Program that minimize the risk of genetic contamination.

Genetic Monitoring Program through *Molecular characterization* helps to determine the breeding behavior of species, individual reproductive success and the existence of gene flow, that is, the movement of alleles within and between populations of the same or related species and its consequences (Papa and Gepts, 2003). Previously, Restriction fragment length polymorphism (RFLPs) has been the choice for many species to measure genetic diversity (Bonstein et al., 1980). But this technique is slow, cumbersome and it requires a large amount of DNA sample. So the use of SNPs (Single nucleotide polymorphism) has replaced the need of RFLP but they are less informative as less allele are present and use a large amount of DNA samples and data analysis (provide reference).

So overcoming the hurdles of RFLP and SNP, the more efficient technique used is microsatellite which are the arrays of repeat sequences that display length variations and different alleles containing different number of repeat units. Microsatellite marker is locus specific, highly polymorphic, co-dominant (Brown, 2001), amenable to analysis by PCR as PCR typing is much quicker and more accurate due to high heterozygosities and very high mutation rates of 10^3 and 10^4 (Jeffreys et al., 1988; Kelly and Mckinnon, 1991; Henderson and Peter, 1992). Thus, microsatellite markers are appropriate for the study of molecular taxonomy, evolution and population genetics (Bowcock *et al.*, 1994) and also applied for DNA typing for individual identification and for assessing the degree of genetic relatedness between individuals. The data interpretation of microsatellite assays is also straightforward and based on visual identification of the polymorphisms. Thus, we implemented microsatellite marker based genetic monitoring program to evaluate the genetic diversity within and between the lines of BALB/c strains after 20th generation of full-sib mating.

Materials and methods

Selection of inbred strain and tail tissue sampling

Four lines of BALB/c strain were selected for the present study that belonged to the CSIR-Indian Institute of Integrative Medicine, Jammu, (CSIR-IIIM). All the breeding mice utilized in the present study were maintained as breeder stock. A total of 80 mice were sampled for tail samples as 10 individuals per lines with equal representation of each sex were chosen randomly as shown in Table 1.

DNA Extraction and Selection of Microsatellite Markers

The tail samples were subjected to DNA extraction by using standard protocol of Sambrook and Russel (1989). In the present study, a set of 14 microsatellite markers were used. Some of these markers selected from mouse genome

informatics (MGI) and were already used in different studies to determine the genetic diversity of mice as shown in Table 2. The primers were synthesized from Sigma.

PCR Amplification

After the extraction of DNA, the microsatellite loci were amplified through polymerase chain reaction (PCR) using primer set for selected microsatellite markers (Table 2). A total of 25µL reaction mixture was prepared for each reaction which included 12.5µl of GoTaq® Green Master Mix, 1µl of 2X upstream primer (10µM), 1µl of downstream primer (10µM), 2µl of DNA template (50ng/µl), 6.5µl of Nuclease-Free Water was used for the thermocycling of the PCR reactions. Initial denaturation at 95 °C for 5 minutes, and then for each of 35 cycles following procedure was used, denaturation at 94 °C for 0.45 seconds, annealing at primer specific temperature for 0.30 second, 72°C for 30 minute for extension process that followed by a final extension at 72 °C for 10 minutes. Using Promega's GoTaq™ DNA Polymerase with Green GoTaq™ Reaction Buffer, eliminated the need to add loading dye to the PCR sample before electrophoresis. PCR product for each sample was loaded along with 50 bp standard ladder onto the 2% agarose gel and allowed to run on the gel at 80 volts for 2 hours.

Statistical Analysis

Each PCR product was genotyped and gel documentation data was analyzed to work out the standard parameters of genetic diversity among lines of BALB/c strain under study. For statistical analysis of genetic diversity studies have done by GenALEX 6.5 (2015) population genetics software (Smouse et al, 2015). This software can be accessed easily online which facilitated to work using Microsoft excel sheet. This way one can easily analyze the numbers of alleles, Shannon information, heterozygosity and genetic distance (Nei) among different population (Shannon C, 1948; Sherwin et al., 2006 & 2010).

Table 1:- Sample (different lines of BALB/c strains) collection from CSIR-IIIM Animal House

Strains	Type	Selected traits	Filial Generation	Institute	No. of female	No. of male	Total sample
BALB/c line-1	Inbred (full-sib mating)	Litter size	>20	CSIR-IIIM	10	10	20
BALB/c line-2	Inbred (full-sib mating)	High weaning weight	>20	CSIR-IIIM	10	10	20
BALB/c line-3	Inbred (full-sib mating)	Low weaning weight	>20	CSIR-IIIM	10	10	20
BALB/c control	Inbred (full-sib mating)	No selection	>20	CSIR-IIIM	10	10	20
					40	40	80

Table 2: - Primer sequences and other information of the microsatellite markers used.

Primer	Repeat Motif	Genetic location	Sequence (5'-3')	(%)GC Content	Fragment Range (bp)	Reference
D1Mit16	(CA)23	1	F: AGAGTTAGCTGCCTAGCTTGAGTG	50	162-197	Dietrich et al. (1992)
			R: TGGAAAGATCTAGGGTTGTCAAAA	37.5		
D1Mit17	(GT)14	1	F: GTGTCTGCCTTTGCACCTTT	50	170-190	Dietrich et al. (1992)
			R: CTGCTGTCTTTCCATCCACA	50		
D1Mit136	(GT)19	1	F: TAGCCCTACACACTGTAGAAATGC	45.8	86-108	Gill and Boyle (2005)
			R: TGAACACAAAAGTAGTAAATGCGTG	37.5		
D1Mit171	(CA)12	1	F: TGCAGATTCAGTCTGCCTTG	50	148-200	Oharaseki et al. (2005)
			R: AGCCATGGGAACACTCTCAC	55		
D1Mit356	(GT)30	1	F: GGGAGAACCTGTCAAGACCA	55	112-152	Namiki et al (2003)
			R: TTTTGGAAATGAGTGTCTGGC	42.8		
D2Mit75	(CA)24	2	F: TCAGCATGTGGATGAATACACA	40.9	16-112	Kaerlsson et al. (2003)
			R: AACTTTTAAAAACTACGAGCGTG	33.3		
D3Mit200	(GT)23	3	F: CAACTTCAGTTTCTCAITTTGAATTG	32	99-131	Cook et al. (2001)
			R: GCAAATGGAAGAGGTTTCTCC	47.6		
D7Mit259	(CA)22	7	F: CCCCTCCTCTGACCTCTT	63.1	116-152	Cattanach et al. (2000)
			R: GTCTCCATGGGAACCACACT	55		
D11Mit227	(CA)34	11	F: CCAGCATTGAACCTGATT	45	116-188	Johannesn et al. (2006)
			R: AAACCCATAGCCTGCATCTG	50		
D11Mit260	(CA)19	11	F: ACTTIGCCTTTATACTATATGGTGG	36	74-118	Farber et al. (2007)
			R: CATTGTTTAGTCTCAGCACCA	39.1		
D13Mit130	(GT)24	13	F: TCTGCTGAAGGCCAGGAC	61.1	139-148	Juriloff et al. (1996)
			R: TTGAAGTGCATGTTGATTTTAATG	28		
D16Mit5	(GT)21	16	F: CGGGGATCATCCCTAAAAAC	50	132-161	Narita et al. (2002)
			R: TCCCAATTCCTCTGTGTC	50		
D17Mit62	-	17	F: CCACATCTTCTAATCCTGTCTCA	45.4	158-176	Gregorova et al. (1996)
			R: CATATAGCCTGAGACATTCTGCC	47.8		
D17Mit124	(CA)15	17	F: TGTTGATGAGATCTTAAATCAGCC	37.5	149-163	Rocha et al. (2004)
			R: TTAACTAGTTGTTATTGCATGTGTG	30.7		

Results

Our Genetic Monitoring Program is grounded in the identification of points in the production program where there is a higher risk of an undetected accidental mating of different lines. By focusing genetic diversity analysis methods on these critical high-risk points, while scrutinizing quality control throughout a genetic monitoring program can prevent a single mistake from becoming catastrophic. At IIIM, Jammu

we used strict strategies at three levels for accurate genetic monitoring program:

Level 1. Visual Phenotypic characteristics of strains.

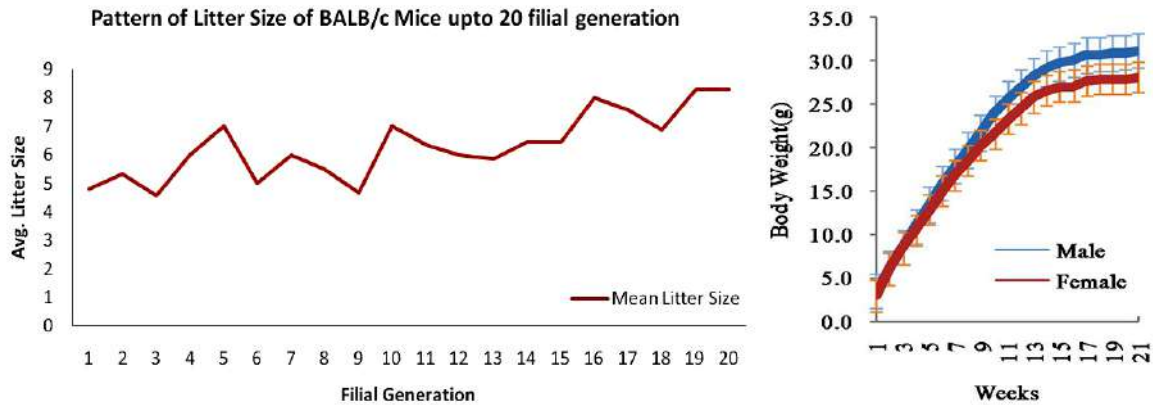
Level 2. Careful record keeping, line segregation and strict pedigree record maintenance.

Level 3. Molecular characterization and Genetic diversity analysis by using Microsatellite Markers.

Results of all the three levels of Genetic Monitoring Program

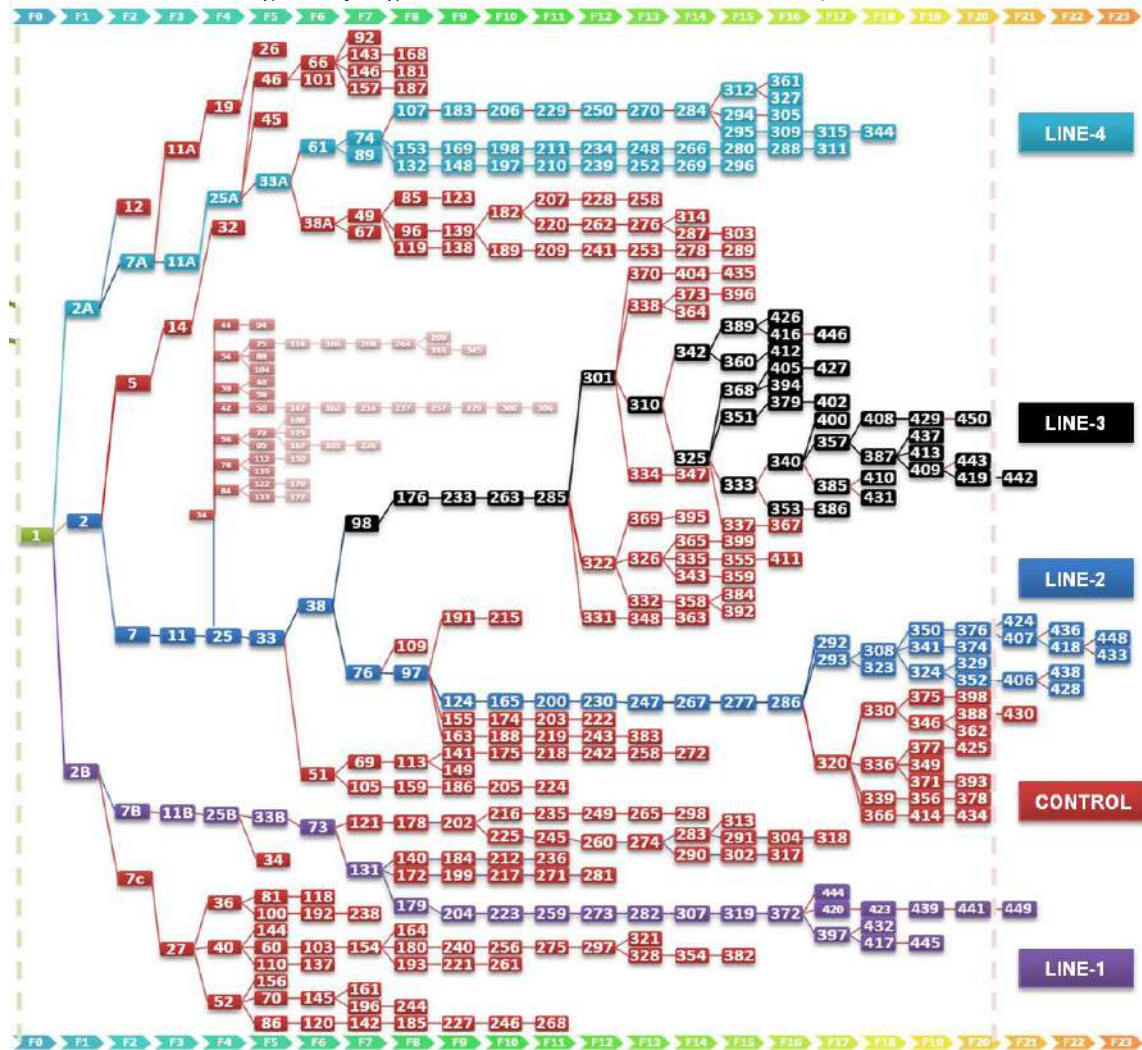
1. Visual characteristics of BALB/c strain (coat color, litter size, weaning weight, growth curve)

Figure-1: Mean litter size and mean growth curve of BALB/c strain at CSIR-IIIM, Jammu.



2. Production procedures that minimize co-localization and/or movement of lines of BALB/c Strain and that promote careful record keeping, line segregation and pedigree maintenance.

Figure-2: pedigree chart of BALB/c strain at CSIR-IIIM, Jammu.



3. Genetic monitoring program outlining 14 different microsatellite markers for 4 different lines of BALB/c mice strain revealed genetic diversity within and between lines of BALB/c mice.

Figure-3: Allelic patterns of male & female BALB/c strain on 2% agarose gel

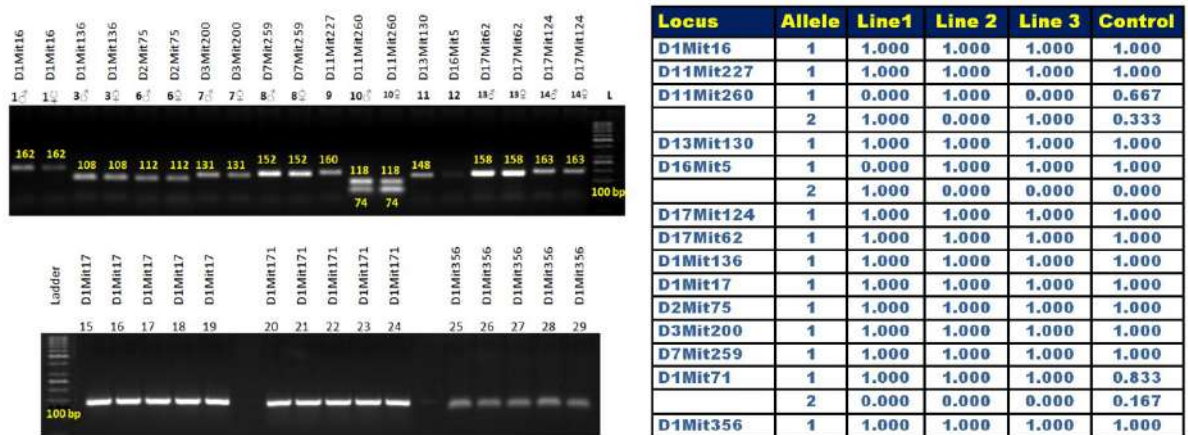
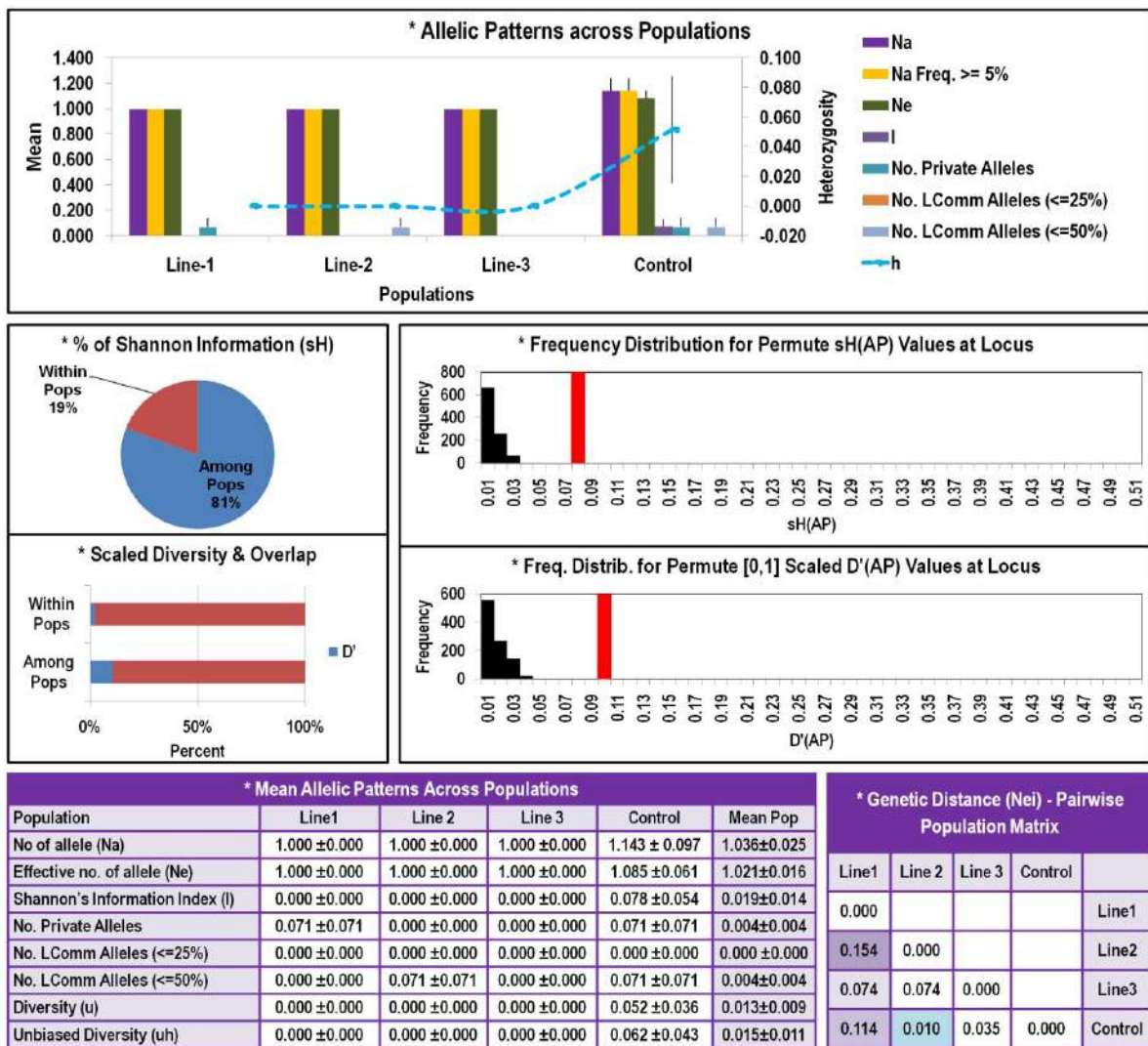


Figure-4: Genetic diversity analysis of different lines of BALB/c strain by GenAIEx 6.5 population genetics software



Discussion

All line mice were genotyped by using established 14 pair of microsatellite markers and genetic diversity analysis (Figure 3). The Effective number of alleles was reported 1.021 ± 0.016 in overall experimental population. Shannon's Index for within lines was 0 and among lines was 0.019 ± 0.014 . The Estimated Diversity (u) was 0 for within lines, 0.052 ± 0.036 for control lines and 0.013 ± 0.009 for among population. Polymorphic information content (PIC) observed in D11Mit260. Line-1 was showing very little Genetic distance with Line-2 (0.154) followed by control line (0.114) and Line-3 (0.074). Control line and Line-2 showed very close genetic distance (0.010). It is concluded that BALB/c strain has no diversity within line and has very little diversity between lines after 20th generation of full-sib mating.

Conclusion

Our experimental results would be easy to obtain using GenAEx 6.5 (2015) population genetics software, which simplifies the genetic diversity analysis from the direct gel

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