

A data format description for genomic breeding evaluation

This document describes the required file format for exchanging data with the National Research Institute of Animal Production.

Strona | 1

Type and format of files required for data exchange

Type of files required for data exchange:

- 1. The file contains genotype information.
- 2. The file contains pedigree information.

Lab ISO17025 requirements

In case the Parentage verification service is obligatory in data verification processes, please be aware that from 2022 onwards, for ICAR accreditation to be granted to laboratories carrying out SNP genotyping, ISO17025 accreditation will be a mandatory minimum requirement as per guidelines. https://www.icar.org/index.php/certifications/certification-and-accreditation-of-dna-genetic-laboratories/guidelines-for-str-and-snp-based-parentage-testing-in-cattle/

Data compression

The genotype file should be compressed with xz format. Compression of the xz file can be performed with software available at http://www.7-zip.org/ for Windows and at http://tukaani.org/xz/ for Linux. If using xz is not possible, gzip compression should be performed.

1. Genotype file

Data from laboratories should be delivered in *Illumina* FinalReport format. Each column in the data set should be <u>tab-delimited</u>:

- 1. SNP-name (according to Illumina nomenclature) SNP Name,
- 2. Short number of individual (14 characters) or International ID of ANIMAL (Interbull ID 19 characters) Sample ID,
- 3. Allele1 (in A/B format) Allele1 AB,
- 4. Allele2 (in A/B format) Allele2 AB,
- Allele1 (w TOP format) Allele1 TOP,
- 6. Allele2 (w TOP format) Allele2 TOP,
- 7. (optional) GC statistic result GC Score,
- 8. (optional) GT statistic result GT Score,

Example:



33/01/2020 1:01 PM Processing Date BovineSNP50_v3_A1.bpm Content Num SNPs 53218 Total SNPs 53218 Num Samples 1526 Total Samples 1526 [data] GC Score SNP Name Sample ID Allele1 - AB Allele2 - AB GT Score 0.9140 ARS-BFGL-BAC-10172 PL000123456789 0.8767 ARS-BFGL-BAC-1020 PL000123456789 A 0.9288 0.8919 В ARS-BFGL-BAC-10245 PL000123456789 0.7227 0.7447 В В ARS-BFGL-BAC-10345 PL000123456789 0 0

Remarks:

- <u>unknown genotypes</u> of SNPs should be marked as "-" according to the default settings of Illumina GenomeStudio or Beeline,
- data exchanged should come from analysis with a call rate of at least 0.95 per individual
- for each type of separate microarray, a file should be created
- animals genotyped using one kind of microarray should be saved into one file
- there is a possibility to replace data in A/B format with a TOP format
- the additional column can be sent in a file with genotype data according to principles of file generation in Illumina GenomeStudio or Beeline,
- please report every exception from the remarks mentioned above before the data transfer!

The naming of the files:

The exchanged file should be named according to the following nomenclature criteria: YYYYMMDD_ NameOfChip_Final_Report.txt.xz

Where:

- YYYYMMDD file preparation date: year (YYYY), month (MM) and day (DD)
- NameOfChip the type of microarray used for genotyping. Possible values are: BovineSNP50v1, BovineSNP50v2, BovineSNP50v3, EuroG10Kv3, EuroG10Kv4, EuroG10Kv5, EuroG10Kv6, EuroG10Kv7, EuroG10Kv8, EuroG10Kv8b, GeneSeekHDv1, GeneSeekHDv1, GeneSeekHDv3, HD, LDv1, LDv1.1, LDv2, ZoetisMD2, GeneSeekGGPv2, GeneSeekGGPv3, GeneSeekGGPv4, Axiom_VM2, Neogen GGP100Kv1, EuroGMDv1, EuroGMDv2, EuroGMDv3, EuroGMDv4, EuroGMDv4.1, EuroGMDv5
- Final_Report permanent unit corresponding to the type of prepared file, whether in Illumina GenomeStudio or Beeline
- extension csv.xz representing xz compressed text file.

^{*}Microarrays not used in Poland are marked with grey colour.



Strona | 2

Example:

20140310_EuroG10Kv3_Final_Report.txt

- indicates data prepared on 10th March 2014 with EuroG10Kv3 microarray

Example:

20170829_BovineSNP50v3_Final_Report.txt

Strona | 3

- indicates data prepared on 29th August 2017 with Illumina BovineSNP50 version v3

2. Pedigree information

Pedigree information can be sent in two formats:

2.1. According to standard Interbull - in format file200 (Table 1).

Table 1. Pedigree file format in file200 type.

Position of the first character in the line	Description	Data type and number of characters	Example	
1	Type of file	character 3	200	
International ID of ANI	MAL			
5	Animal breed ¹	character 3	HOL	
8	Animal first registration country ²	character 3	CAN	
11	Sex ³	character 1	М	
12	Animal ID ⁴	character 12	000000A12345	
International ID of Sire of ANIMAL				
25	Sire breed	character 3	HOL	
28	Country of first registration of sire ²	character 3	CAN	
31	Sex	character 1	М	
32	Sire ID ⁴	character 12	556912367589	
International ID of Dan	n of ANIMAL			
45	Dam breed	character 3	HOL	
48	Country of first registration of dam ²	character 3	CAN	
51	Sex	character 1	F	
52	Dam ID ⁴	character 12	123569874521	
65	Animal birthdate (YYYYMMDD)	integer 8	19870215	
74	Status of animal ⁵	integer 2	10	
77	Birthdate of first AI daughters (YYYYMMDD)	integer 8	19890314	
86	Name of the animal	character 30	Cantarello	
The national ID of the a	animal			
117	Animal breed ¹	character 3	HOL	
120	Country of first registration of the animal ²	character 3	CAN	
123	Sex ³	character 1	M	
124	Animal ID ⁴	character 12	000000A12345	



137 Country of data exchange	character 3	CAN
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- *1 HOL= Holstein Friesian breed (HF) for black and white and red and white type; BSW= Brown Swiss; JER= Jersey; SIM= Simental; RED= national red breeds.
- *2 Country where the animal was first registered, described according to ISO alpha-3 country designation table (for example, POL for Poland, DEU for Germany, etc.)
- *3 Sex M –sire, bull, F cow, dam, heifer

Strona | 4

- *4 Unique, <u>12-digit number of animal</u>, all shorter names (below 12 characters) should be completed with "zero" starting from the beginning of ID (from the left side)
- *5 Bull status: 00 unknown, also used for cows, 10 AI bull, 20 other bull
 - Data should be provided as a text file (*.txt)

 - Lack of birth date (YYYYMMDD) of the animal should be marked as 00000000 (8x0)
 - The absence of the birth date of the first daughter AI (YYYYMMDD) should be marked as 00000000 (8x0)
 - Gaps in the bull status should be marked as 00 (2x0)
 - The total record line should contain 140 characters, including spaces

Example:

INFORMATION OF PEDIGREE IN FILE200 FORMAT SHOULD BE PREPARED ACCORDING TO RULE AS DEEP AS POSSIBLE, for both the sire and dam side.

2.2. The data format of animal identification (Table 2).

Table 2. The data format of animals sent to microarray genotype analysis.

Columns	number of characters	Format	Remarks
Sample number	20		
Chip ¹	14		BovineSNP50 EuroG10K
Sample type	8		blood semen ear tissue other
Date of sampling	8	YYYY-MM-DD	
Number of the animal (Short name)	14		

¹ BovineSNP50 for the animals that do not require imputation; EuroG10K for the animals that require imputation



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^{*}spaces marked as _

National Institute of Animal Production – data exchange, 2024-08-28

Name of the animal	30			
Breed/type	2			
Birthdate	8	YYYY-MM-DD		
Sex	1		M - bull F - cow	
Embryotransfer	2		ET	ona 5
Number of sire (Short number)	14			·
Name of sire	30			
Number of dam (Short number)	14			
Name of dam	30			
Number of sire dam (Short number)	14			
Name of sire dam (Short number)	30			
Genotype operator ¹	16		WCHiRZ Poznań MCHiRZ Łowicz MCB Krasne SHiUZ Bydgoszcz PFHBiPM Warszawa	

^{*1} or other institution authorized for that purpose.

FILE SHOULD BE SAVED IN CSV (comma-separated values) https://pl.wikipedia.org/wiki/CSV (file format).

The naming of file with pedigree

File with pedigree information should be subjected to the following naming criteria: YYYYMMDD_pedigree_file200.txt.xz YYYYMMDD_pedigree_dane.csv.xz

Where:

- YYYYMMDD data of file preparation containing year (YYYY), month (MM), and day
 (DD) of file preparation, the date should correspond to the date of file with genotypes
- pedigree constant unit corresponding to the type of prepared file
- file200.txt.xz constant unit corresponding to the type of prepared pedigree file of animals in file200 – file200 format, compressed in xz format
- dane.csv.xz constant unit corresponding to the type of animal identification data data compressed with xz format.

Example:

20140310_pedigree_file200.txt.xz

- indicates compressed pedigree data (xz) in file200 prepared on 10th March 2014.

Example:

20170829_pedigree_dane.csv.xz

- indicates compressed animal identification data (xz) in csv file prepared on 29th August 2017.



Genotype file transfer to ftp server

According to individually sent information about data transfer, prepared files should be verified and compressed before being sent to ftp server.

Pedigree file transfer to ftp server and email

Strona | 6

According to individually sent information about data transfer, prepared files should be verified and compressed before being sent to ftp server.

Data with pedigree information both in file200 format and/or animal identification file format should be sent to Kacper Żukowski at kacper.zukowski@iz.edu.pl and Monika Skarwecka monika.skarwecka@iz.edu.pl.

Contact information



National Research Institute of Animal Production 1, Krakowska Street PL32083 Balice Poland

The contact person regarding genotype data transfer is Kacper Żukowski, email: kacper.zukowski@iz.edu.pl.

The contact person regarding the pedigree information transfer is Monika Skarwecka, email: monika.skarwecka@iz.edu.pl.

Format history

2024-08-28	-	chips added
2023-10-30	-	added the TOP format on a par with AB, @ changed
2021-05-06	-	added the TOP format on a par with AB, @ changed
2019-12-21	-	format revision, chips added
2020-06-07	-	add ISO requirements
2019-05-30	-	format revision, chips added
2018-01-15	-	added English version of a document
2017-10-09	-	addition of file 200 to format description
2017-09-29	-	systematization of file format

