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Maximum number of total born piglets in a parity and individual ranges in litter size expressed as specific characteristics of sows

Gertraude Freyer

Abstract

Background: The objective of this study was to underline that litter size as a key trait of sows needs new parameters to be evaluated and to target an individual optimum. Large individual variation in litter size affects both production and piglet's survival and health negatively. Therefore, two new traits were suggested and analyzed. Two data sets on 5509 purebred German Landrace sows and 3926 Large White and crossing sows including at least two parental generations and at least five parities were subjected to variance components analysis.

Results: The new traits for evaluating litter size were derived from the individual numbers of total born piglets (TBP) per parity: In most cases, sows reach their maximum litter size in their fourth parity. Therefore, data from at least five parities were included. The first observable maximum and minimum of TBP, and the individual variation expressed by the range were targeted. Maximum of TBP being an observable trait in pig breeding and management yielded clearly higher heritability estimates ($h^2 \sim 0.3$) than those estimates predominantly reported so far. Maximum TBP gets closer to the genetic capacity for litter size than other litter traits. Minimum of TBP is positively correlated with the range of TBP ($r_p = 0.48$, $r_g > 0.6$). The correlation between maximum of TBP and its individually reached frequency was negative in both data sets ($r_p = -0.28$ and -0.22, respectively). Estimated heritability coefficients for the range of TBP comprised a span of $h^2 = 0.06$ to 0.10.

Conclusion: An optimum both for maximum and range of total born piglets in selecting sows is a way contributing to homogenous litters in order to improving the animal-related conditions both for piglets' welfare and economic management in pig.

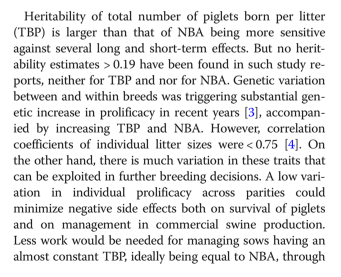
Keywords: Pig, Litter size, Individual maximum, Variation, Heritability, Trait correlation

Background

In pig breeding, the number of piglets born alive per parity (NBA) is a trait of economically importance, but affected by multiple factors [1]. Heritability is always reported to be relatively low, and the wise breeding decision is an enduring challenge. Many biologic effects on litter size emphasize the problem: healthy conditions of the sow, specific effects due to the birth process, individual maternal influences. Multifold environmental impacts do play a role as well. One specific example on intrinsic effects is the content of serum immunoglobulin being lower in piglets from larger litters [2].

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all their parities. Nursing and feeding conditions could be stabilized. A reduced individual variation in TBP would help to improving efficiency of piglet production and animal welfare. Targeting an individual optimum in litter size is presumably a thorny issue, but worth investigating.

The objective of this study was on the genetic capacity of TBP expressed by the individual maximum TBP (Max_TBP) and on the individual variation in TBP (Range_TBP). Two data sets based on genetically different pools from commercial swine production in Germany were subjected to analyzing phenotypic trait observations by means of a simple but widely used and recognized methodical approach in order to estimate heritability coefficients for these new parameters for litter traits.

Material and Methods

Phenotypic data

Fecundity data collected in 30 farms from January 1997 through June 2010 was provided by the Hybridschweinezuchtverband Nord/Ost e.V. Malchin, Germany. In the first data set, 15,271 German Landrace sows (RA01) stemming from 1425 sires belonged to about 800 contemporarily farrowing groups. Heritability coefficients on this large dataset based on first parities have already been reported ($h^2 = 0.10$ and 0.13 for NBA respective TBP) [5]. In the present study, 5509 sows with at least five parities were considered (Table 1). Most phenotypic

Table 1 Basic statistics on the investigated traits individualmaximum and minimum of number total born piglets (Max_TBPand Min_TBP), and individual range of number total born piglets(Range_TBP), number of total born piglets and number of pigletsborn alive in first parity (TBP_first and NBA_first) in GermanLandrace sows (RA01) and Large White sows including crossingsows (RACOM) with at least five individual parities each

	Overall observed							
	Mean	Standard deviation	Minimum	Maximum				
German Landrace sows (R	A01, n = 5	5509)						
Min_TBP	8.67	2.581	1	17				
Max_TBP	15.46	2.181	9	29				
Range_TBP	7.21	2.908	1	22				
TBP_first	11.46	2.850	1	21				
NBA_first	10.81	2.739	1	20				
Overall parity number	6.57	1.480	5	15				
Large White and crossing	sows (RA	COM, n = 392	26)					
Min_TBP	9.34	2.700	1	17				
Max_TBP	15.63	2.260	8	26				
Range_TBP	6.29	2.803	1	22				
TBP_first	11.53	2.905	2	20				
NBA_first	10.84	2.779	1	20				
Overall parity number	6.90	1.830	5	16				

individual maximum total born piglets of sows occurred in their fourth parities (Fig. 1). This suggests that the true individual potential denoted by Max TBP mostly is observable when the sow has finished at least four parities. The pedigree contained 30,620 individuals in total including 6168 base animals and two or more parental generations of the recorded sows in RA01. The second data set (RACOM) meeting the same conditions as above comprised 3926 German Large White purebred sows and sows from crossings with German Landrace, stemming from 436 sires (details on crossing sows: dam was Large White for 211 sows, sire was Large White for 2053 sows). Therefore, three breed groups had to be allowed for in RACOM. The whole pedigree was larger in RACOM, containing 37,003 individuals with 6567 base animals. Inbreeding was not an issue, neither in RA01 and nor in RACOM.

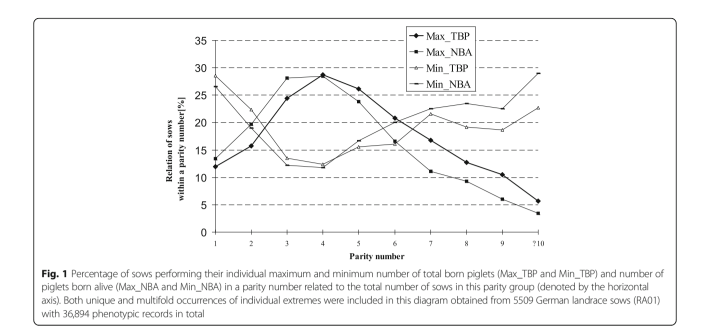
Description of investigated traits and influencing factors

The target trait was the individual maximum of TBP (Max_TBP). As a secondary trait, a characteristic for describing individual variation in TBP of sows should be easily observable in swine production was on focus. Range_TBP was used here on the basis of individual Max_TBP and individual minimum TBP (Min_TBP). A sufficient number of individual parities of each sow **h** was necessary to calculate

Range_TBP_{hk} = $Max_TBP_{hi} - Min_TBP_{hj}$,

where \mathbf{i} and \mathbf{j} denoted different parity numbers yielding the single maximum respective minimum of TBP and \mathbf{k} was the number of individually available parities.

Min_TBP and Max_TBP were in fact new traits based on their first individual occurrence. Range_TBP describes a specific situation that cannot be expressed by original measurements. Calculating Range_TBP is therefore comparable to the blood pressure amplitude in medicine. Minimum and maximum were obtained from different parities of a single sow. E.g., the specific parity number and contemporarily group of sows (expressed by herd, year and season of parity) leading to the specific trait observation was considered as fixed effect on Max_TBP and Min_TBP. A peculiarity was that the year of first farrowing had a significant effect on Max_ TBP (P < 0.0001) suggesting a targeted selection effect. In Range_TBP, the situation was different in terms that significance of fixed effects was found for herd, year and season related to Min_TBP. Further, the number of available parities (in analyses of using greater than five parities) resulted in a significant effect on Range_TBP, but not the parity number related to Min_TBP or Max_TBP.



All significant effects were allowed for trait specifically in statistical analyses. For all targeted traits, skewness and kurtosis were negligibly small. The precondition of a normal distribution was fulfilled.

NBA at first parity (NBA_first) has apparently been the most respected trait for production purposes so far as suggested from the majority of relevant reports. Therefore, it was contemplated in parallel throughout the study.

Statistical analyses

Genetic parameters of the investigated traits were estimated on the basis of the animal model

$$\mathbf{y} = \mathbf{X}\mathbf{b} + \mathbf{Z}\mathbf{u} + \mathbf{e},$$

where **y** is the vector of trait observations (phenotypes), **b** vector of fixed effects (breed code, farm, herd-group, year and season of parity, actual parity number (see also above for further significant effects)), **u** vector of direct additive genetic (animal) effects, and **e** the vector of residual effects. **X** denotes the incidence matrix of fixed effects, **Z** matrix of direct additive-genetic effects. **u** and **e** were normally distributed (0, $\mathbf{A} \otimes \mathbf{G}$) with numerator relationship matrix **A** and additive genetic variance-covariance structure **G**, and (0, $\mathbf{I}_e \otimes \mathbf{R}$) for **e**, respectively, with. \mathbf{I}_e identity matrix and **R** variance-covariance structure of residuals. The symbol \otimes denotes the Kronecker product.

The variance components analyses aiming at estimating heritability coefficients and their standard errors (SE) were carried out by means of univariate modeling and based on one individual trait observation/trait value each for a single sow. Computer programs VCE 5.1.2 and PEST V4.0 were used for the estimating heritability (and SE) based on REML and were run on a LINUX machine [6, 7]. SAS package [SAS Institute Inc., Cary, NC, US, versions 9.2 and 9.4] was used for general statistics including Pearson correlation coefficients (accompanied by the error probability P) and for testing fixed effects and group differences by means of GLM and MIXED procedures.

Min_TBP and Max_TBP were considered once as unique trait values at their first occurrence. A repeatability model was therefore not applied for any of the traits.

Results

Heritability estimates

The relative additive variance components representing the heritability coefficients are shown for both samples (Table 2). In fact, results of two analyses per sample are listed: (i) results from analyses based on all parities \geq five, and (ii) results based on exactly five parities. Heritability estimates of Max_TBP were within a span of 0.29 to 0.39. Standard errors were low in both samples (SE = 0.02 ... 0. 04). Heritability estimates for Min_TBP were considerably lower, namely $h^2 = 0.07 \dots 0.11$ (in most cases, SE < 0.02).

Heritability estimates of Range_TBP were lowest in RA01 ($h^2 = 0.06 \pm 0.01$) compared to RACOM ($h^2 = 0.10 \pm 0.02$), whereas the use of exactly five parities let to closer results between both samples, but accompanied by larger SE (Table 2).

Frequency of individual maxima and minima in contemplation to Range_TBP

About 26 percent of sows reached their Max_TBP more than once during their available parities. Three to four percent yielded the same maximum more than twice. The results were almost equal for RA01 and RACOM.

Table 2 Estimates of relative additive genetic variance
components expressing the heritability coefficients (h^2) and their
standard errors (VC _{rel} \pm SE) on the investigated traits ^a Min_TBP,
Max_TBP and Range_TBP and in addition on NBA in first parity
based on univariate analyses

Trait		Estimated variance components in relation to the total phenotypic variance					
	Additive genetic	Residual					
RA01 (German Land	race sows, <i>n</i> = 5509)						
(i) information on	five or more parities used	for estimation					
Min_TBP	0.112 ± 0.014	0.888 ± 0.014					
Max_TBP	0.298 ± 0.021	0.702 ± 0.021					
Range_TBP	0.056 ± 0.011	0.944 ± 0.011					
NBA_first	0.123 ± 0.017	0.877 ± 0.017					
(ii) information or	n exactly five parities used f	or estimation					
Min_TBP	0.070 ± 0.015	0.930 ± 0.015					
Max_TBP	0.283 ± 0.023	0.717 ± 0.023					
Range_TBP	0.066 ± 0.012	9.934 ± 0.012					
RACOM (Large Whit	te and crossing sows, $n = 39$	926)					
(i) information on	five or more used for estin	nation					
Min_TBP	0.098 ± 0.018	0.902 ± 0.018					
Max_TBP	0.373 ± 0.030	0.627 ± 0.030					
Range_TBP	0.104 ± 0.022	0.896 ± 0.022					
NBA_ first	0.116 ± 0.020	0.884 ± 0.020					
(ii) information or	n exactly five parities used f	or estimation					
Min_TBP	0.070 ± 0.024	0.930 ± 0.024					
Max_TBP	0.390 ± 0.040	0.610 ± 0.040					
Range_TBP	0.084 ± 0.026	0.916 ± 0.026					
^a investigated traits we	ere individual maximum and r	minimum of number total					

^ainvestigated traits were individual maximum and minimum of number total born piglets (Max_TBP, Min_TBP), and individual variability in number total born piglets (Range_TBP); number of piglets born alive in first parity (NBA_first) for comparing to heritability estimates from earlier studies

The relation was less for Min_TBP. 20 percent of sows showed their Min_TBP twice or more.

Negative phenotypic correlation coefficients being very similar in both samples were estimated for the individual frequency of Max_TBP and Range_TBP ($r_p = -0.25$ and -0.23, P < 0.0001). The even stronger negative correlation of individual frequency of Min_TBP and Range_TBP was almost the same in RA01 and RACOM ($r_p = -0.32$ and -0.31, P < 0.0001).

In most cases of RA01, multiple maxima occurred for Max_TBP = 13 ... 16 piglets (in 996 cases, respective 69 percent of sows with multiple maxima). In RACOM, the "preferred span of maxima" was Max_TBP = 13 ... 17 piglets. Multiple minima were observed most frequently for Min_TBP = 9 ... 11 in RA01 (in 56 percent of sows with multiple minima), and Min_TBP = 10 ... 12 in RACOM. In total, RACOM was more homogenous in Range_TBP, confirmed by lower mean and lower standard deviation (Table 1).

Phenotypic and genetic trait correlations

Almost all estimated trait phenotypic correlation coefficients were significant (Table 3). The estimated genetic correlation coefficients were highly positive for Max_TBP and Range_TBP: $r_g = 0.617 \pm 0.061$ in RA01 and $r_g = 0.761 \pm 0.067$ in RACOM. The genetic correlation of Range_TBP and Min_TBP were not significant, $r_g = 0.141 \pm 0.132$ in RA01 and $r_g = 0.214 \pm 0.180$ in RACOM.

Phenotypic correlation coefficients for piglets born alive in first parity (NBA_first) and Range_TBP were significantly negative ($r_p = -0.15$ in RA01 and $r_p = -0.22$ in RACOM, Table 3). This seems advantageous when focusing on a very early breeding decision. However, positive genetic correlations of NBA_first and Range_TBP minimize the expectations ($r_g = 0.33$ in RA01 and $r_g = 0.31$ in RACOM, SE = 0.09 and 0.10, respectively).

Discussion

In commercial swine production, litter size is a key trait. However, individual numbers of total born and piglets born alive vary in subsequent parities. So far, this fact has hardly been targeted as a special parameter of sows. In this study, individual ranges of TBP were chosen to do so as a simple trait in order to draw attention to this problem and to suggest coping with.

Selection on larger TBP increases NBA but in many cases it also increases the number of still born piglets. For this reason and to respecting the individual feeding capacity of a sow, searching for an individual optimum TBP should be a target in pig breeding. In the investigated material, a clear trend of increasing mean performances in TBP and NBA of first parity were observed during 1997 to 2008. At the same time, there was no increase in mean Range_TBP, but standard deviations of Range_TBP showed an increased trend from 1997 to 2008 very clearly (not shown in further detail). This suggests potential for respecting individual variation without necessarily decreasing litter sizes on average.

Respecting individual Max_TBP and individual Range_ TBP could be a way for optimizing the management process in swine production and improving animal welfare directly on the sow's level. Litter size traits in pigs are likely more affected by maternal than by paternal components. One might argue that the number of five individual parities to evaluate the genetic capacity of TBP and individual variation is a challenging limitation. The preference on fast selection decisions in sows focuses on their performances in NBA_first.

Which parameter really matters for evaluating a sow for litter size with respect to efficiency? Observed Max_TBP is an individual parameter for the genetic capacity of litter size as clearly suggested by larger heritability estimates compared to traditionally used litter size traits (Table 2). **Table 3** Phenotypic correlation coefficients of individual variability in number total born piglets (Range_TBP), maximum and minimum number of total born piglets (TBP_Max and TBP_Min), and in addition both the number piglets born alive in first parity (NBA_first) and the individual relative number of still born piglets (SB_rel)^a obtained from RA01 (n = 5509, above diagonal) and from RACOM (n = 3926, below diagonal), Pearson correlation coefficients were significant at P < 0.0001, besides those marked by superscript letters

	RA01 (above dia	RA01 (above diagonal)									
		Range_TBP	Max_TBP	Min_TBP	NBA_first	SB_rel					
RACOM	Range_TBP		0.478	-0.669	- 0.145	0.079					
(below diagonal)	Max_TBP	0.475		0.316	0.277	0.287					
	Min_TBP	-0.672	0.332		0.392	0.159					
	NBA_first	-0.219	0.257	0.452		0.011 ^b					
	SB_rel	0.086	0.286	0.148	-0.003 ^c						

^aSB_rel was based on phenotypic on all available parities observations on still births adjusted for fixed effects of breed indicator (within RACOM), parity number, herd, year and season using the basic data set (in total, 63,000 records in RA01, 41,500 records in RACOM) for all sows included in the current study ^bP = 0.4256

 $^{c}P = 0.8437$

Targeting an optimum TBP does not mean selecting for highest Max_TBP. This trait should be evaluated in order to find a basis for choosing the optimum. Simultaneously, a small individual Range_TBP should be preferred in future breeding objectives. Comparing single sows yielding extremely different Range_TBP reveals the practical value of low individual variation in litter size. Sows with a Max_ TBP of 13 to 16 showed the lowest Range_TBP and simultaneously the most repeatable Max_TBP, accompanied by fewest still births. The significant connection between still births and Range_TBP is also shown by dividing sows into range classes and by phenotypic correlations (Fig. 2, Table 3) and by means of eight single sows being extremely different in their Range_TBP (Table 4).

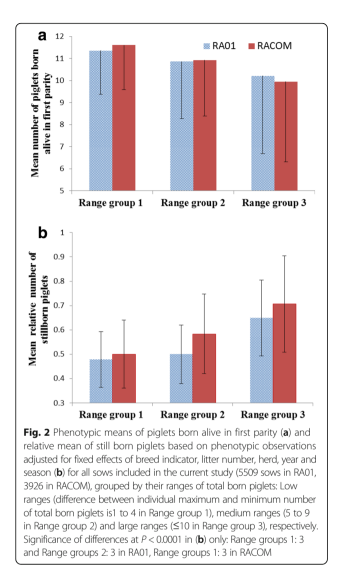
In the present study, the relative genetic variance (heritability) estimated for Range_TBP was about six to 10 percent. RACOM was more homogenous in Range_TBP than RA01. Due to the large part of crossing sows included in RACOM, this could be a response of favorable non-additive components triggering more stabilizing factors on expressing these traits. On the other hand, despite fewer sows with trait observations, the pedigree was much larger for RACOM than for RA01, a fact that could have caused higher heritability estimates.

Selecting sows with an individual optimum TBP respective NBA for breeding could contribute to more phenotypically (and especially environmentally) balanced sustainability. Priority should be given to feeding capacity and maternal behavior.

Birth weight of each single piglet could likely be an additional co-variable in evaluating litter size traits [8]. However, such data was not available for this study.

Table 4 Single sows from RACOM with exactly five parities and extremely low or extremely large ranges of their number in total born piglets per parity (Range_TBP), maximum number of total born piglets (Max_TBP) marked in bold, and the related absolute number of still born piglets (SB)

		Parity number											
group with respect to Range_TBP and breed		1		2		3		4		5		Total	
	ID	TBP	SB	TBP	SB	TBP	SB	TBP	SB	TBP	SB	TBP	SB
Low Range_TBP													
Crossing sow	196679	16	2	15	0	16	0	15	0	15	0	77	2
Large White	285278	13	2	13	1	13	1	14	1	13	0	66	5
Large White	106820	11	0	12	0	12	0	12	0	11	0	58	0
Crossing sow	200791	10	0	10	1	10	0	11	1	11	0	52	2
Large Range_TBP													
Large White	106324	17	5	20	2	18	6	7	0	5	0	67	13
Crossing sow	194908	10	0	17	2	18	0	17	4	1	0	63	6
Large White	107757	19	0	18	1	17	1	4	0	13	0	71	2
Large White	272177	9	0	11	0	14	2	26	13	14	1	74	16



NBA in first parity was analyzed in parallel resulting in very similar heritability estimates as reported before [5]. A selection effect on sows under study was therefore not suspected.

No earlier reports based on individual maxima and on individual female variation of TBP have been found in the literature related to pig breeding. A recent study carried out in the Netherlands was based on boars' observations through their daughters [9]. For a deeper insight into the genetics of litter size and related traits, SNP technology is being widely adapted. Such studies have been reviewed [10], and candidate genes of reproduction traits in sows were reported based on different pig populations, partly with contradictory results [11–13].

Non-additive effects of single candidate genes (or chromosomal segments) interacting with others could play a different role for TBP and NBA in first and higher parities [11]. Therefore, patterns of age-dependently (inter-) acting genes, and genes affecting environmental sensitivity, could also affect the individual variation of traits related to litter size. Two candidate genes contributing to variation in TBP were reported [8]. One of them is involved in buffering environmental and genetic factors. From the same study, a genetic correlation between TBP and its variation based on boars' breeding values was reported ($r_g = 0$. 49 [8]). This was very similar to the phenotypic correlation of Range_TBP and Max_TBP in the present study ($r_p = 0.48$, Table 3).

The results of the study reported here suggest that an individual optimum number of total born and alive born piglets per parity of a sow can be found by respecting individual Max_TBP and Range_TBP. Heritability of sows' individual Max_TBP is higher than heritability estimates for litter size traits as published so far. More detailed investigations on the basis of a genome wide association study and surely using methods allowing for genotype by environmental interactions could lead to increasing the knowledge on the responsible genes, multifold induced interactions and their functions. Focusing on sows being extremely different in their individual variation in litter size would be of value in this continuing research process.

Conclusions

Using the individual maximum number of total born piglets as a new trait in genetic analyses reveals considerably higher heritability estimates ($h^2 \sim 0.3$) than those from using ordinary litter size data repeatedly reported before. Information on the individual maximum is more suited to reflect the genetic proliferative potential of the sow. Many sows reached their maximum of 13 to 16 piglets in a parity more than once. Therefore, this trait is suggested to find an individual optimum litter size for improving both the management process and animal's health. A sufficient evaluation of sows regarding both their individual capacity in litter size and the related individual variation (e.g. range as a simple secondary trait) is possible if data of at least five parities is available. Individual range of litter size in sows has a heritable component ($h^2 = 0.06$ to 0.10).

Abbreviations

Max_NBA: Individual maximum number of piglets born alive related to a specific parity; Max TBP: Individual maximum number of total born piglets of a sow related to a specific parity; Min_NBA: Individual minimum number of piglets born alive related to a specific parity; Min_TBP: Individual minimum number of total born piglets of a sow related to a specific parity; NBA: Individual number of piglets born alive per parity; NBA_first: Individual number of piglets born alive in first parity; P: Error probability; RA01: Data set from German Landrace; RACOM: Data set from Large White and crossing sows; Range TBP: Individual range of number of total born piglets of a sow (individual variation); SB: Number of still born piglets of a sow; SB_rel: Individual number of still born piglets in a parity adjusted for fixed effects of herd, year and season of farrowing; SE: Standard error; TBP: Total number of piglets per parity; VCE_{rel}: Relative variance component; Further, symbols on population genetic parameters were used: h^{2} - heritability coefficient; r_a and r_p - genetic and phenotypic correlation coefficients. Other specific symbols were used only once in the methods section (with explanation)

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Authors' contributions

The author read and approved the final manuscript.

Ethics approval

Not applicable.

Competing interests

The author declares that she has no competing interests.

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