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Comparison between the three porcine RN genotypes for growth, carcass composition and meat quality traits

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Abstract — A three-step experimental design has been carried out to add evidence about the existence of the RN gene, with two segregating alleles RN\textsuperscript{−} and rn\textsuperscript{+}, having major effects on meat quality in pigs, to estimate its effects on production traits and to map the RN locus. In the present article, the experimental population and sampling procedures are described and discussed, and effects of the three RN genotypes on growth and carcass traits are presented. The RN genotype had no major effect on growth performance and killing out percentage. Variables pertaining to carcass tissue composition showed that the RN\textsuperscript{−} allele is associated with leaner carcasses (about 1 s.d. effect without dominance for back fat thickness, 0.5 s.d. effect with dominance for weights of joints). Muscle glycolytic potential (GP) was considerably higher in RN\textsuperscript{−} carriers, with a maximum of a 6.85 s.d. effect for the live longissimus muscle GP. Physico-chemical characteristics of meat were also influenced by the RN genotype in a dominant way, ultimate pH differing by about 2 s.d. between homozygous genotypes and meat colour by about 1 s.d. Technological quality was also affected, with a 1 s.d.

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decrease in technological yield for RN\textsuperscript{−} carriers. The RN genotype had a more limited effect on eating quality. On the whole, the identity between the acid meat condition and the RN\textsuperscript{−} allele effect is clearly demonstrated (higher muscle GP, lower ultimate pH, paler meat and lower protein content), and the unfavourable relationship between GP and carcass lean to fat ratio is confirmed.

**Résumé** — Comparaison des trois gènes RN chez le porc pour les caractères de croissance, de composition de la carcasse et de qualité de la viande. Un protocole expérimental en trois étapes a été mis en œuvre pour confirmer l’existence du gène RN, avec deux allèles en ségrégation RN\textsuperscript{−} et rn\textsuperscript{+}, à effet majeur sur la qualité de la viande chez le porc, en estimer les effets sur les caractères de production et en déterminer la localisation génétique. Dans cet article, la population expérimentale et les procédures d’échantillonnage sont décrites et discutées, puis les effets des trois génotypes RN sur les caractères de croissance et carcasse sont présentés. Le gène RN n’a pas d’effet notable sur les performances de croissance et le rendement de carcasse. Les variables relatives à la composition tissulaire de la carcasse indiquent que l’allèle RN\textsuperscript{−} est associé à des carcasses plus maigres (environ 1 écarts type sans dominance pour les épaisseurs de lard dorsal, 0,5 e.t. avec dominance pour les poids de morceaux). Le potentiel glycolytique musculaire (GP) est beaucoup plus élevé chez les porteurs de RN\textsuperscript{−}, avec un écart maximum de 6,85 e.t. pour la mesure in vivo du GP sur le muscle longissimus. Les caractéristiques physico-chimiques de la viande sont également influencées par le gène RN d’une façon non additive, le pH ultime différant d’environ 2 e.t. entre homozygotes et la couleur de la viande d’environ 1 e.t. La qualité technologique est aussi affectée, avec 1 e.t. de diminution du rendement technologique chez les porteurs de RN\textsuperscript{−}. Le gène RN au locus RN a un effet plus limité sur les qualités sensorielles de la viande. Globalement, l’identité entre les caractéristiques de la viande acide et les effets de l’allèle RN\textsuperscript{−} est clairement démontrée (potentiel glycolytique musculaire supérieur, pH ultime inférieur, viande plus pâle, concentration en protéines inférieure) et la relation défavorable entre GP et rapport muscle/gras est confirmée.

**1. INTRODUCTION**

Pigs showing an abnormally large extent of post mortem muscle pH fall were first described by Monin and Sellier [26] as characteristic of the Hampshire breed (i.e. “Hampshire effect”). In 1986, Naveau [28] postulated the existence of a single major gene to explain the occurrence of this “acid meat” condition in two composite lines, Penshire and Laconie, built from Hampshire blood at a rate of 1/2 and 1/3, respectively. In the latter study, the genetic determination of an indicator of the technological yield of cured-cooked ham processing, the “Napole yield” (RTN: Rendement Technologique Napole [29]), was explored. The postulated major gene was called RN, the dominant allele responsible for the decrease of RTN being RN\textsuperscript{−} and the normal recessive allele being rn\textsuperscript{+}. This hypothesis was further confirmed by Le Roy et al. [20] using segregation analysis methods on RTN field data. Moreover, Wassmuth et al. [35], analysing Hampshire crossbred populations, demonstrated the segregation of a major gene (denoted HF for “Hampshirefaktor”) influencing meat quality in the same way as RN. However, all these results were obtained from a posteriori statistical analyses of field data and had to be confirmed using an experimental design specifically devoted to the evaluation of RN gene effects.
It was early postulated that the “Hampshire effect” arises from higher muscle glycolytic potential (GP) [11, 26]. That the primary effect of the RN$^-$ allele is to strongly increase GP was a logical and attractive hypothesis. Several studies have therefore consisted of comparing animals of either high GP or low GP, within Hampshire crossbred populations, in order to estimate the effects of the RN$^-$ allele [7-10, 23, 24, 30]. However, this classification based on GP is not fully satisfying because (1) the RN gene was initially found through its effect on RTN, and the effect of the RN$^-$ allele on GP has never been properly demonstrated, (2) only RN$^-$ carriers and non-carriers have been compared instead of the three genotypes RN$^-$/RN$^-$, RN$^-$/rn$^+$ and rn$^+$/rn$^+$, and (3) estimates of the RN$^-$ effect could be biased due to the selection procedure which led to comparison of animals with extreme GP phenotypes and thus potentially extreme values for correlated traits.

A three-step experimental design has been implemented to add evidence about the existence of the RN gene [21], to estimate its effects on various traits while avoiding the above-mentioned drawbacks, and to map the RN locus [25]. The aim of the present article is: (1) to describe the experimental population; (2) to give elements for validation of the comparison between RN genotypes; (3) to report the effects of the three RN genotypes on the three main traits characterising the Hampshire effect and the acid meat condition (RTN, GP and ultimate pH), as well as on growth performance and carcass quality. Results concerning the effects of the three RN genotypes on chemical composition, enzyme activities and myofiber characteristics of muscle are reported elsewhere [19].

2. MATERIALS AND METHODS

2.1. Experimental design

2.1.1. General principles

The experiment was carried out on Le Magneraud INRA Unit (Surgères, Charente Maritime, France). Founder animals were from the Laconie composite line, created in 1973 and selected by the Pen ar Lan breeding company (Maxent, Ille et Vilaine, France). This line was originally founded with Hampshire, Piétrain and Large White blood in equal proportions. The present design was primarily constructed to compare the three RN genotypes and was set up according to three principles: (1) comparisons had to be made between individuals differing by their RN genotype but sharing similar polygenic background; (2) the RN genotype had to be determined using the initial definition of the gene, i.e. its effect on the RTN trait; and (3) the effects of the RN genotype had to be measured on animals of a priori known genotypes, i.e. animals born from proven homozygous parents.

The design comprised three steps: (1) animals supposed to be heterozygous were intercrossed to produce a segregating population of RN$^-$/RN$^-$, RN$^-$/rn$^+$ and rn$^+$/rn$^+$ individuals sharing similar polygenic background; (2) males and females from this segregating population were progeny tested with the aim of determining their RN genotype; (3) offspring from proven homozygous parents were produced in a “diagonal” cross for comparing the three RN genotypes.
2.1.2. Herd foundation

Prior to the start of this experiment, RTN had been recorded on 9726 Laconie animals (from 156 sires and 937 dams) and all corresponding breeding boars and sows were genotyped for RN from analysing RTN records of their progeny. Simplified segregation analysis as described by Elsen and Le Roy [6] was used assuming segregation of the two alleles RN\(^{-}\) and rn\(^{+}\) in both sexes. Boars and sows having an estimated probability of 1 to be homozygous (either rn\(^{+}\)/rn\(^{+}\) or RN\(^{-}\)/RN\(^{-}\)) were chosen to establish the experimental population. The consistency of predicted genotypes of parents, mates and grand parents was checked prior to the final choice. Five females classified as RN\(^{-}\)/RN\(^{-}\) and 4 females classified as rn\(^{+}\)/rn\(^{+}\) were mated to 6 males classified as rn\(^{+}\)/rn\(^{+}\), and pregnant sows were transferred to Le Magneraud where they farrowed. Two groups of piglets from the resulting litters were considered: (1) a group of animals born from rn\(^{+}\)/rn\(^{+}\) dams, assumed to be homozygous rn\(^{+}\)/rn\(^{+}\), and among which 4 males and 8 females were used to found a tester line (T); (2) a group of animals born from RN\(^{-}\)/RN\(^{-}\) dams, assumed to be heterozygous RN\(^{-}\)/rn\(^{+}\), and among which 6 males and 19 females were used to found the segregant population (S).

2.1.3. Progeny test

These 6 sires and 19 dams gave birth to 273 candidate offspring among which RN\(^{-}\)/RN\(^{-}\), RN\(^{-}\)/rn\(^{+}\) and rn\(^{+}\)/rn\(^{+}\) were expected in proportions 1/4, 1/2 and 1/4, respectively. Due to limited experimental facilities, a small part of these candidates could be progeny tested for RTN. In order to avoid a random loss of homozygotes, preselection of the animals to be progeny-tested was performed on the basis of an individual in vivo measurement of muscle GP (IVGP) at 70 kg live weight [34]. Thus, among 67 boars and 83 gilts measured for IVGP, 16 and 43 were kept for being submitted to the progeny test, 6 and 12 with low IVGP (lower than 200 \(\mu\)mol·g\(^{-1}\), a priori rn\(^{+}\)/rn\(^{+}\)) and 10 and 31 with high IVGP (greater than 300 \(\mu\)mol·g\(^{-1}\), a priori RN\(^{-}\)/RN\(^{-}\) or RN\(^{-}\)/rn\(^{+}\)). The T line, supposed to be homozygous recessive rn\(^{+}\)/rn\(^{+}\), consisted of 6 sires and 34 dams. In order to verify the RN genotype of these animals, a progeny test was also implemented, with each T dam giving one litter sired by a T boar. A segregation analysis was performed on the progeny-test RTN data [21] to estimate the posterior genotype probabilities of all sires and dams (Fig. 1). Results showed that one T boar was certainly heterozygous. As a consequence, the litters sired by this boar were deleted from the design, and only 37 of the 43 females from the S population were validly tested. From both groups of S animals classified as homozygous (RN\(^{-}\)/RN\(^{-}\) or rn\(^{+}\)/rn\(^{+}\)), 3 boars and 11 sows were kept to generate the animals of the third step.

2.1.4. Diallel cross

The 22 sows were distributed in three 3-week-spaced farrowing batches. One of the rn\(^{+}\)/rn\(^{+}\) dams gave no litter, 7 dams (5 rn\(^{+}\)/rn\(^{+}\) and 2 RN\(^{-}\)/RN\(^{-}\)) gave only one litter, and the 14 others gave 2 litters, with alternate genotypes for 10 of them, i.e. one heterozygous litter and one homozygous litter. Finally, 12, 11 and 12 litters were produced in the RN\(^{-}\)/RN\(^{-}\), RN\(^{-}\)/rn\(^{+}\) and rn\(^{+}\)/rn\(^{+}\)
genotypes, respectively and it was possible to balance the distribution of RN genotypes within each slaughter series. Numbers of pigs recorded for each group of traits are given by RN genotype in Table I.

**Table I.** Numbers of pigs recorded for each group of traits.

| Trait                                      | RN genotype |
|--------------------------------------------|-------------|
|                                            | RN⁻/RN⁻ | RN⁻/rn⁺ | rn⁺/rn⁺ |
| Postweaning growth performance (¹)         | 103       | 92       | 69       |
| In vivo muscle glycolytic potential        | 98        | 88       | 66       |
| Carcass composition, Napole yield and physico-chemical muscle characteristics | 90        | 73       | 57       |
| Loin eye area, pH₁, post mortem glycolytic potential and cured-cooked ham processing ability | 37        | 38       | 39       |
| Eating quality of meat                     | 17        | 17       | 17       |

(¹) In brackets, numbers of pens.

**Figure 1.** Results of the progeny test for RTN: relationships of RN genotype estimated by segregation analysis with family mean, within family standard deviation and own IVGP value (in white, parents with IVGP greater than 300 μmol·g⁻¹; in black, parents with IVGP smaller than 200 μmol·g⁻¹).
2.2. Traits

2.2.1. Growth performance

Piglets were weaned at 28 days of age and moved to the fattening building at 77 days. They were penned in groups of 6 to 12 animals, each pen including females or castrated males from the same RN genotype. During the fattening period, animals were fed ad libitum a standard pelleted diet (crude protein: 17.0%; crude fat: 1.5%; crude fiber: 4.5%; ash: 6.8%; lysine: 0.85%; ME: 3091 kcal/kg⁻¹). Average daily gain was recorded individually from 30 to 100 kg live weight. Food conversion ratio from 30 to 100 kg live weight was calculated on a pen basis as the ratio of feed consumed to live weight gain.

2.2.2. Live muscle glycolytic potential

A shot-biopsy sample of longissimus lumborum muscle was taken at 71 ± 7 kg live weight, as described by Talmant et al. [34]. Biopsy samples were immediately trimmed of skin and fat, and homogenised in 10 mL of 0.55 M perchloric acid. At the laboratory (Station de recherches sur la viande, INRA, Theix, France), 0.5 mL of the homogenate was used for simultaneous determination of glycogen, glucose-6-phosphate and glucose [5]. The rest of the homogenate was centrifuged at 2500 × g during 10 min, and the supernatant was used for lactate determination [2]. Muscle GP, in μmol equivalent lactate per g of fresh tissue, was calculated according to Monin and Sellier [26]: GP = 2([glycogen] + [glucose-6-phosphate] + [glucose]) + [lactate]. The sum of glycogen, glucose-6-phosphate and glucose concentrations will be referred to as “glycogen concentration” in the following.

2.2.3. Carcass composition

Pigs were slaughtered at 107 ± 9 kg live weight in a commercial abattoir (Celles sur Belle, Charente Maritime, France). On the day after slaughter, the carcass (with head, feet and leaf fat) was weighed, and killing out percentage was calculated as the ratio of cold carcass weight to live weight. Carcass length (from the first cervical vertebra to the anterior edge of the pubial symphysis) and midline back fat thickness (at the shoulder, back and rump levels) were measured on the right side of the carcass. Then, this side was weighed and divided into seven joints (ham, loin, shoulder, belly, back fat, leaf fat and feet) according to a standardised cutting method [1]. Weights of joints were recorded and carcass lean percentage (CLP) was estimated according to the following equation (1): CLP = −42.035 + (1.282 ham weight + 1.818 loin weight + 0.616 shoulder weight + 0.701 belly weight + 0.040 leaf fat weight − 0.678 back fat weight) / half carcass weight. Carcass compactness was defined as the ratio of loin weight to carcass length. Loin eye area was measured at the last rib level by planimetry using a tablet digitizer (Hitachi).

2.2.4. Physico-chemical characteristics of muscle

At 35 min after slaughter, a sample of longissimus muscle was removed from the right half-carcass at the last rib level and homogenised in 18 mL of 5 mM iodoacetate for pH measurement (pH1). At the same time, samples of three
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muscles, differing in their metabolic and contractile properties (longissimus, semimembranosus and semispinalis capitis) [16,27], were taken for determination of post mortem glycogen concentration, lactate concentration and GP (PMGP), as previously described.

The day after slaughter, the following traits were recorded on loins and hams:

- pH24 of adductor femoris, biceps femoris, gluteus superficialis, longissimus, semimembranosus and semispinalis capitis muscles. Measurements were made directly on muscles using a combined glass electrode (Ingold, Mettler Toledo, Switzerland) and a portable pH meter (CG818, Schott Geräte, Germany);

- colour (L*, a* and b* values) of biceps femoris, gluteus superficialis and longissimus muscles, using a Minolta chromameter CR-300;

- water-holding capacity of biceps femoris, gluteus superficialis and longissimus muscles, as assessed by the “filter paper imbibition time” method [3], i.e. the time required for complete wetting of a 1 cm² filter paper piece put on the freshly cut surface of the muscle.

2.2.5. Technological meat quality

The “Napole” curing-cooking yield was recorded on a 100 g sample of semimembranosus muscle. The method used was that described by Naveau et al. [29] except that the muscle sample was removed from the right half-carcass the day after slaughter and not on the slaughter line. However, the time of meat maturation at 4 °C, about 24 h, remained the same. One ham was processed into cured-cooked ham by the Eden company (La Chataigneraie, Vendée, France). Raw weight (X1), deboned-defatted weight (X2), weight after curing (X3) and weight after cooking (X4) were recorded in the course of processing. The following yields were calculated: anatomic yield (X2/X1), curing yield (X3/X2), cooking yield (X4/X3), technological yield (X4/X2) and overall yield (X4/X1).

2.2.6. Eating quality

The day after slaughter, three slices (1 cm thick) were removed from the loin at the last rib level, vacuum-packed and stored at −20 °C for about six months. Then, the frozen samples were thawed at 4 °C for 24 h, deboned and cooked on an electric grill for 4 min at 170 °C. In a total of 17 testing sessions, grilled chops were scored by a taste panel of 12 trained people for the following traits: visual compactness at cutting, tenderness, juiciness, mellowness and pork flavour intensity. Each descriptor was scored on a 10-point scale, from zero (very low) to 10 (very high).

2.3. Statistical methods

2.3.1. Validation of prediction and comparison of the RN genotypes

In the course of the experiment, progeny tested animals from the segregant and tester populations have been selected considering their estimated RN genotype obtained from simple two-generation segregation analyses of RTN records, as described by Le Roy et al. [21]. Few errors were detected in the expected rn⁺/rn⁺ genotyping of tester animals, suggesting possible misclassifications in founders. Considering all pedigree and RTN information collected in the design as a whole should improve the accuracy of RN genotype prediction.
A second source of bias is inevitably expected from the selection of homozygous parents of the diallel cross: these animals were selected as extreme for the RN phenotype of their progeny test offspring, which should increase the differences in polygenic means between RN\(^{-}/RN^{-}\) and rn\(^{+}/rn^{+}\) selected parents. Analysing the genotypic effect of the diallel step animals without taking into account these phenomena could give an overestimation of the RN gene effects on RTN and correlated traits.

Guo and Thompson [13] proposed a pedigree analysis method which considers genealogy and performance records from the whole pedigree and thus makes a full use of available information for a single trait. The main feature of this method is the joint use of an EM algorithm and the Gibbs sampler for estimating the parameters of the mixed model of inheritance (major gene + polygenes). A more accurate genotyping of individuals can be expected from such a pedigree analysis as compared to the two-generation approach. Moreover, when records used for selection of parents are included in the analysis, a less biased estimation of parameters should be obtained, as far as the results found by Henderson [14] and others can be generalised to the mixed inheritance context.

The estimates of RN genotype effects on RTN were estimated from three approaches. The reference was the pedigree analysis with all RTN records described above. To evaluate the potential bias due to both genotype misclassification and selection of parents of the diallel cross, the two following simplified analyses were performed: a full pedigree analysis with the only diallel step RTN records; a classical mixed model (fixed + random effects), where the same genealogical information was used, but where the RTN of the last generation only was considered and RN genotypes were supposed to be known without error. The second approach did not consider the selection problem, the third approach did not neither consider the selection nor the misclassification problems. The complete pedigree starting from the founder animals chosen in Maxent comprised 1791 animals among which 1641 had a RTN record. All these data were considered in the reference pedigree analysis whereas only records of the 220 individuals of the diallel step were considered in the two simplified approaches. It was expected that, if little difference is found, the classical mixed model approach could provide a reliable estimates of the RN effects on all traits measured.

The Guo and Thompson [13] algorithm has been implemented in Fortran language with the following characteristics chosen after a number of trials: a dememorisation step of 100 Gibbs samples; 500 EM steps; a Monte Carlo sample size of 100; 20 Gibbs samples between two consecutive Monte Carlo samplings. More than \(10^6\) samples have thus been generated. In order to increase mixing, the proposition of Janss et al. [15] for sampling of major genotypes has been retained: Gibbs sampling has been applied to the subvector of parents + final progenies (not having offspring) rather than to all individuals independently.

Three fixed effects have been included in the model, in accordance of their statistical significance in preliminary analyses of variance: sex (2 levels: female and castrated male), HAL genotype, determined using molecular genotyping [4] (2 levels: NN and Nn), and date of slaughter (107 levels). For any individual, the probability of each of the three RN genotypes was estimated by the mean, computed during the last EM step (100 samples), of this RN genotype
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2.3.2. Estimation of RN genotype effects

Classical maximum likelihood analysis was performed using the PEST software [12]. Starting from the final generation of pigs, i.e. those recorded in the diallel step, pedigree was followed back up to the founders in order to constitute the pedigree file which contained 340 animals over 6 generations. The inbreeding option was used.

Traits were analysed in univariate models. The RN genotype of recorded individuals was supposed to be perfectly known and was considered as a fixed effect (3 levels: RN$^-$/RN$^-$, RN$^-$/rn$^+$ and rn$^+$/rn$^+$). As stated above, three other fixed effects were included in the model: sex (2 levels), HAL genotype (2 levels) and environmental effect, i.e. date of biopsy for muscle GP (6 levels), fattening batch for growth and carcass composition traits (6 levels) and date of slaughter for meat quality traits (11 levels). Initial weight for average daily gain, live weight at biopsy for GP and live weight at slaughter for carcass and meat quality traits, were included as covariates. Litter effect and additive genetic value were considered as random effects. The corresponding variance components ($\sigma^2_c$ and $\sigma^2_a$, respectively) could not be estimated from the present data due to the small size of data sets, and they were derived from average values of heritability ($h^2$) and common litter environment ($c^2$) reported in the literature [32]. The phenotypic variance $\sigma^2_p$ of each trait was estimated using the GLM procedure of SAS [31] and was set equal to the residual mean square of a fixed model analysis of variance including the same effects as those contained in the above-mentioned mixed models. Variance components were defined as $\sigma^2_c = c^2 \sigma^2_p$, $\sigma^2_a = h^2 \sigma^2_p$ and $\sigma^2_e = \sigma^2_p - \sigma^2_c - \sigma^2_a$.

3. RESULTS AND DISCUSSION

3.1. Validation of RN genotypes comparison

Table II reports the predicted RN genotypes of progeny-tested animals using either full pedigree analysis or two-generation segregation analysis. In both approaches, a parent has been given a genotype G if the estimated probability of G was higher than 0.80. When none of the three possible genotypes had a probability higher than 0.80, the genotype was considered as unknown (denoted “??”).

With this threshold, few discrepancies were found between the two genotyping methods. One progeny-tested male was classified as RN$^-$/rn$^+$ with the two-generation segregation analysis and as rn$^+$/rn$^+$ with the pedigree analysis. The latter classification is consistent with his own low (179 $\mu$mol·g$^{-1}$) in vivo GP (not considered in the analyses). Regarding sows, three discrepancies were observed (1 RN$^-$/rn$^+$ changed to RN$^-$/RN$^-$ and 2 rn$^+$/rn$^+$ from the tester line changed to RN$^-$/rn$^+$), without any clear explanation, except the fact that...
they had a limited number of offspring (23, 6 and 11). Thirteen undetermined animals were more clearly genotyped with the pedigree approach. It should be emphasised that none of the boars and sows used as parents of the diallel-step offspring or of the resource families for linkage analyses showed a change in RN genotype in this retrospective study.

Based on the full pedigree approach, the RTN means were 83.2, 83.6 and 91.0% for $RN^-/RN^-$, $RN^-/rn^+$ and $rn^+/rn^+$ animals respectively, with a within-genotype standard deviation of 2.8. These figures confirm that the RN major gene is a dominant gene with a difference of 2.8 standard deviation (s.d.) units between means of homozygotes, an estimate very close to that found in the original study of Le Roy et al. [20] (2.9 s.d. units in the Laconie line). The within-major genotype heritability estimate was 0.46 in the present data set, to be compared with the estimate of 0.28 found by Le Roy et al. [20]. This increase in heritability is consistent with the expected better control of environment in the present experiment.

When the full pedigree approach was applied limiting the RTN information to the diallel step, the genotype means for RTN (in %) were 82.2, 83.3 and 91.2 for $RN^-/RN^-$, $RN^-/rn^+$ and $rn^+/rn^+$ animals respectively. Based on the second simplified approach (classical animal model), the contrasts between genotype means for RTN, (in %) were estimated as $-8.2 \pm 0.8$ and $-7.8 \pm 0.6$ for $RN^-/RN^- - rn^+/rn^+$ and $RN^-/rn^+ - rn^+/rn^+$, respectively, using the variance component estimates from the pedigree analysis ($\sigma_p = 2.8; h^2 = 0.46$). A bias, reaching about 5%, was then probably due to the selection of parents of the diallel step, the estimates being close to those previously found [20]. Then, the diallel-step could be considered as a random sampling of RTN polygenes, allowing to estimate the RN gene effect on other recorded traits with a bias lower than 5%.

In the following comparisons, the PEST software was used and both litter and additive genetic random effects were taken into account in the model of analysis, genetic parameters being set to classically accepted values. With that method, the same two contrasts between genotype means for RTN were estimated as $-8.4 \pm 0.7$ and $-7.8 \pm 0.6$ with a within-genotype standard deviation being equal to 2.6 and $h^2$ and $c^2$ coefficients being set to 0.30 and 0.05, respectively. Several tests showed that the estimates of RN genotype means for RTN are quite robust to variation in parameters $h^2$ and $c^2$.  

### Table II. Distribution of breeding boars and sows according to their RN genotype as determined by either segregation analysis or pedigree analysis.

| Genotype predicted from pedigree analysis | Genotype predicted from two-generation segregation analysis | Total |
|----------------------------------------|-------------------------------------------------|-------|
| $RN^-/RN^-$                           | $RN^-/rn^+$                                     | 19    |
| $RN^-/rn^+$                           | $RN^-/rn^+$                                     | 19    |
| $rn^+/rn^+$                           | $rn^+/rn^+$                                     | 44    |
| ?                                     | ?                                               | 1     |
3.2. Estimation of RN genotype effects

Tables III to VII give results of the RN genotype comparison. Only contrasts between genotypic means can be estimated without bias, and results are presented relative to the control rn+/rn+ genotype ($\mu_{rn^-/rn^-} - \mu_{rn+/rn+}$ and $\mu_{rn^-/rn+} - \mu_{rn+/rn+}$ contrasts). Least squares means for the rn+/rn+ genotype ($\mu_{rn+/rn+}$), and the within-genotype standard deviations ($\sigma_p$), as computed by the SAS GLM procedure, are also given. For each trait, both tests of significance of the RN genotype effect (test of the $\mu_{rn^-/rn^-} - \mu_{rn+/rn+} = 0$ hypothesis) and of the dominance effect (test of the $d = 0$ hypothesis, with $d = \mu_{rn^-/rn+} - 0.5(\mu_{rn^-/rn^-} + \mu_{rn+/rn+})$) are shown.

3.2.1. Growth performance

Estimated effects of the RN genotype on growth traits (Tab. III) did not significantly differ from 0, except for average daily gain. For this trait, the heterozygote RN$^-$/rn+ had a significant advantage over the two homozygous genotypes which were very close to each other. The dominance effect was highly significant ($P < 0.01$) and was estimated as 42 g day$^{-1}$, i.e. one half of the phenotypic standard deviation of the trait. Such a situation of over dominance is fairly surprising, but it should be mentioned that a favourable effect of the RN$^-$ allele on daily gain was also found by Enfält et al. [7] comparing RN$^-$/rn+ and rn$^+$/rn+ animals.

3.2.2. Carcass composition

Effects of the RN genotype on carcass composition traits are given in Table IV. There was no RN genotype effect on killing out percentage or carcass compactness, but RN$^-$/RN$^-$ animals were longer than RN$^-$/rn+ and rn$^+$/rn+ pigs. These results are agree with those of Enfält et al. [7] and Reinsch et al. [30] which found no difference between RN$^-$/rn+ and rn$^+$/rn+ animals for these traits.

On the whole, variables pertaining to carcass tissue composition showed that the RN$^-$ allele is associated with leaner carcasses. Except for the measurement at the shoulder, back fat thickness was decreased by about 1 s.d. in homozygous carriers RN$^-$/RN$^-$, heterozygotes being intermediate between the two homozygotes and the dominance effect being very close to 0. Concerning the weight of carcass joints, the same trend was observed, with a significant increase in weight of lean joints (ham and loin) and a concomitant, though smaller, decrease of weight of fat joints (belly, back fat and leaf fat). However, the estimated RN effect was lower than for backfat thickness, with differences of only about 0.5 s.d. between means of the two homozygotes. Furthermore, the dominance effect was generally significant, and the heterozygous and homozygous carriers were not different. Consequently, carcass lean content was increased by about 0.75 s.d. in RN$^-$ carriers with a situation of complete dominance. Loin eye area, measured only on a subsample of animals, followed a similar pattern.
Table III. Effect of the RN genotype on growth performance traits ($h^2 = 0.35; c^2 = 0.15$).  

| Trait                              | $\sigma_p^b$ | $\mu_{m+}^b$ | $\mu_{RN^-/RN^-} - \mu_{m+}$ | $\mu_{RN^-/RN^+} - \mu_{m+}$ | $p_1 > \chi^2$ | $d^d$ | $p_2 > \chi^2$ |
|------------------------------------|--------------|--------------|-------------------------------|-------------------------------|---------------|-------|---------------|
| Weight at 21 days of age (kg)      | 1.4          | 6.1±0.2      | 0.3±0.4                       | -0.2±0.4                      | 0.449         | -0.3±0.3 | 0.269         |
| Initial weight (kg)               | 4.5          | 32.8±0.6     | -1.0±1.3                      | 0.1±1.1                       | 0.575         | 0.6±0.9 | 0.478         |
| Final weight (kg)                 | 9.0          | 103.9±1.2    | 1.1±2.6                       | 3.9±2.2                       | 0.163         | 3.3±1.8 | 0.063         |
| Average daily gain (g.day$^{-1}$) | 81           | 879±11       | 8±24                          | 47±20                         | 0.030         | 42±16   | 0.009         |
| Food conversion ratio$^e$         | 0.14         | 2.88±0.05    | -0.12±0.07                    | -0.03±0.06                    | 0.188         | 0.03±0.05 | 0.537         |

$a$ $h^2$ and $c^2$: coefficients of heritability and of common environment, when it was necessary, used for genetic evaluation by PEST.  
$b$ $\sigma_p$ and $\mu_{m+}$: estimates of the within genotype standard deviation and of the within $m+$/m$^+$ genotype mean (± standard error) computed by SAS GLM.  
$c$ $\mu_{RN^-/RN^-} - \mu_{m+}$, $\mu_{RN^-/RN^+} - \mu_{m+}$ and $p_1 > \chi^2$: estimates of the contrasts between genotypic means (± standard error) and level of significance of the RN effect (test of the "$\mu_{RN^-/RN^-} - \mu_{m+} = 0$ and $\mu_{RN^-/RN^+} - \mu_{m+} = 0$" hypothesis) computed by PEST.  
$d$ $d$ and $p_2 > \chi^2$: estimate (± standard error) ($d = \mu_{RN^-/RN^+} - 0.5 (\mu_{RN^-/RN^-} + \mu_{m+})$) and level of significance (test of the "$d = 0$" hypothesis) of the dominance effect computed by PEST.  
$e$ Estimates on a pen basis using SAS GLM.
Table IV. Effect of the RN genotype on carcass composition traits ($h^2 = 0.50$; $c^2 = 0.05^a$).

| Trait                          | $\sigma_p^b$ | $\mu_{RN^+/RN^-}$ | $\mu_{RN^-/RN^+}$ | $p_1 > \chi^2c$ | $d^d$ | $p_2 > \chi^2d$ |
|-------------------------------|-------------|-------------------|-------------------|-----------------|------|---------------|
| Killing out percentage (%)    | 1.5         | 80.6 ± 0.2        | -0.1 ± 0.5        | 0.4 ± 0.4        | 0.307| 0.4 ± 0.3     |
| Carcass length (mm)           | 22          | 958 ± 3           | 15 ± 7            | 1 ± 5            | 0.039| -6 ± 4        |
| Backfat thickness (mm)        |             |                   |                   |                  |      |               |
| Shoulder                      | 4.0         | 34.2 ± 0.6        | -0.7 ± 1.2        | -0.4 ± 0.9       | 0.827| -0.1 ± 0.7    |
| Back                          | 3.3         | 23.8 ± 0.5        | -3.0 ± 1.0        | -1.5 ± 0.8       | 0.009| -0.0 ± 0.6    |
| Rump                          | 2.5         | 14.5 ± 0.4        | -2.6 ± 0.7        | -1.6 ± 0.6       | 0.002| -0.3 ± 0.5    |
| Average                       | 2.5         | 24.2 ± 0.4        | -2.1 ± 0.7        | -1.2 ± 0.6       | 0.017| -0.2 ± 0.5    |
| Weight of joints (kg)         |             |                   |                   |                  |      |               |
| Head                          | 0.29        | 4.05 ± 0.05       | 0.17 ± 0.09       | 0.03 ± 0.07      | 0.089| -0.05 ± 0.05  |
| Ham                           | 0.36        | 10.80 ± 0.06      | 0.13 ± 0.11       | 0.25 ± 0.09      | 0.011| 0.18 ± 0.07   |
| Loin                          | 0.60        | 12.56 ± 0.09      | 0.35 ± 0.18       | 0.42 ± 0.14      | 0.014| 0.24 ± 0.11   |
| Belly                         | 0.32        | 5.09 ± 0.05       | -0.24 ± 0.09      | -0.17 ± 0.08     | 0.030| -0.05 ± 0.06  |
| Shoulder                      | 0.30        | 6.72 ± 0.05       | -0.11 ± 0.09      | -0.11 ± 0.07     | 0.276| -0.05 ± 0.05  |
| Backfat                       | 0.45        | 3.32 ± 0.07       | -0.25 ± 0.13      | -0.24 ± 0.11     | 0.074| -0.11 ± 0.08  |
| Leaf fat                      | 0.11        | 0.55 ± 0.02       | -0.06 ± 0.03      | -0.07 ± 0.03     | 0.052| -0.03 ± 0.02  |
| Feet                          | 0.06        | 0.86 ± 0.01       | 0.04 ± 0.02       | 0.00 ± 0.01      | 0.054| -0.02 ± 0.01  |
| Lean content (%)              | 2.7         | 59.3 ± 0.4        | 1.9 ± 0.8         | 2.2 ± 0.6        | 0.002| 1.3 ± 0.5     |
| Compacity (g/mm$^{-1}$)        | 0.7         | 13.1 ± 0.1        | 0.2 ± 0.7         | 0.4 ± 0.6        | 0.737| 0.3 ± 0.5     |
| Loin eye area (cm²)           | 5.5         | 49.6 ± 0.9        | 2.7 ± 1.8         | 3.5 ± 1.5        | 0.061| 2.1 ± 1.2     |

$^a$ $h^2$ and $c^2$: coefficients of heritability and of common environment, when it was necessary, used for genetic evaluation by PEST.

$^b$ $\sigma_p$ and $\mu_{RN^+/RN^-}$: estimates of the within genotype standard deviation and of the within $RN^+/RN^+$ genotype mean (± standard error) computed by SAS GLM.

$^c$ $\mu_{RN^-/RN^-}$ - $\mu_{RN^+/RN^+}$, $\mu_{RN^+/RN^-}$ - $\mu_{RN^+/RN^+}$ and $p_1 > \chi^2$: estimates of the contrasts between genotypic means (± standard error) and level of significance of the RN effect (test of the $\mu_{RN^-/RN^-} - \mu_{RN^+/RN^+}$ and $\mu_{RN^-/RN^+} - \mu_{RN^+/RN^+} = 0$ hypothesis) computed by PEST.

$^d$ $d$ and $p_2 > \chi^2$: estimate (± standard error) ($d = \mu_{RN^-/RN^+} - 0.5 (\mu_{RN^-/RN^-} + \mu_{RN^+/RN^+})$) and level of significance (test of the “$d = 0$” hypothesis) of the dominance effect computed by PEST.
That carcass lean meat content is higher in RN− carriers than in RN− non-carriers has been consistently reported [7, 23, 30]. However, an effect of the RN gene on backfat thickness was only found in the present study, i.e. the only one including the homozygous RN−/RN− genotype. Dominance relationships for backfat thickness (additivity of alleles) differed from those for lean meat content or weights of joints (complete dominance of RN− over rn+). This could probably be explained by the RN effect on carcass length. Indeed, RN−/RN− animals were longer but with lower backfat thickness when compared with RN−/rn+ animals, which resulted in the same weight of backfat for both carrier genotypes.

The favourable effect of RN− on carcass lean to fat ratio is in accordance with the positive genetic correlation of lean to fat ratio with muscle GP found by Larzul et al. [18] in a presumably RN−-free population. Higher muscle GP appears to be genetically associated with leaner carcass regardless of whether the increase in GP is due to the RN− allele or to polygenes.

### 3.2.3. Muscle glycolytic potential

Results concerning the muscle GP measured in vivo (I.V.) and post mortem (P.M.), are presented in Table V. All variables pertaining to GP and its components, except the PM lactate concentration in the longissimus muscle, were very strongly affected by the RN genotype. In longissimus muscle GP, the difference between homozygotes reached 6.85 s.d. in vivo but “only” 3.45 s.d. post mortem. On the other hand, this difference was lower when the measured muscle was more oxidative, i.e. 3.45 s.d. in the longissimus, 3.09 s.d. in the semimembranosus and 1.09 s.d. in the semispinalis capitis muscle. Regarding the muscle GP, the effect of dominance of RN− over rn+ was significant in the longissimus and semimembranosus muscles and close to significance in the semispinalis capitis muscle. The RN−/rn+ pigs did not significantly differ from the RN−/RN− pigs in GP of the semimembranosus and semispinalis capitis muscle whereas the RN−/rn+ pigs showed significantly lower IVGP and PMGP values than the RN−/RN− pigs in longissimus muscle.

Regarding the components of muscle PMGP, the effect of RN− on residual glycogen was larger in the “white” longissimus and semimembranosus muscles than in the “red” semispinalis capitis muscle. The two “white” muscles somehow differed regarding the effect of RN− on PM lactate concentration: there was no effect of RN− (and even a slightly negative effect) in the longissimus muscle whereas the overall effect of RN− was positive in the semimembranosus muscle.

All these results showed that the primary effect of the RN gene is certainly to increase muscle GP. As stated above, this widely accepted hypothesis had never been properly demonstrated. Here, the RN genotypes being established from RTN measurements, the identity between the “RN gene effect” and the “Hampshire effect” is clearly proven. Furthermore, all previous studies on the RN gene effects did not consider the homozygous genotype RN−/RN− and so could not estimate the effect of dominance. The dominance of the RN− allele for muscle GP can be assumed to be complete in the semimembranosus and semispinalis capitis muscles and almost complete in the longissimus muscle.
Table V. Effect of the RN genotype on muscle glycolytic potential \((h^2 = 0.30; \ c^2 = 0.10)^a\).

| Trait                              | Muscle               | \(\sigma_p^b\) | \(\mu_{rn^+}/rn^+\) | \(\mu_{RN^-}/RN^- - \mu_{rn^+}/rn^+\) | \(\mu_{RN^-}/rn^- - \mu_{rn^+}/rn^+\) | \(p1 > \chi^2^c\) | \(d^d\) | \(p2 > \chi^2^d\) |
|-----------------------------------|----------------------|----------------|---------------------|------------------------------------------|------------------------------------------|-------------------|--------|----------------|
| I.V. Glycolytic potential \((\mu \text{ mol/g})\) | longissimus          | 20             | 167 ± 3             | 137 ± 5                                  | 110 ± 4                                  | 0.000             | 41 ± 4 | 0.000         |
| P.M. Glycolytic potential \((\mu \text{ mol/g})\) | longissimus          | 33             | 110 ± 5             | 114 ± 9                                  | 86 ± 8                                   | 0.000             | 29 ± 7 | 0.000         |
|                                  | semimembranosus      | 32             | 104 ± 5             | 99 ± 9                                   | 87 ± 8                                   | 0.000             | 37 ± 7 | 0.000         |
|                                  | semispinalis capitis | 21             | 44 ± 3              | 23 ± 6                                   | 20 ± 5                                   | 0.000             | 8 ± 4  | 0.072         |
| P.M. Glycogen concentration \((\mu \text{ mol/g})\) | longissimus          | 18             | 35 ± 3              | 59 ± 5                                   | 44 ± 5                                   | 0.000             | 14 ± 4 | 0.000         |
|                                  | semimembranosus      | 16             | 28 ± 3              | 47 ± 5                                   | 36 ± 4                                   | 0.000             | 13 ± 3 | 0.000         |
|                                  | semispinalis capitis | 8              | 5 ± 1               | 8 ± 2                                    | 6 ± 2                                    | 0.002             | 2 ± 2  | 0.246         |
| P.M. Lactate concentration \((\mu \text{ mol/g})\) | longissimus          | 14             | 40 ± 2              | −4 ± 4                                   | −3 ± 3                                   | 0.591             | −1 ± 3 | 0.254         |
|                                  | semimembranosus      | 19             | 47 ± 3              | 6 ± 5                                    | 14 ± 5                                   | 0.013             | 11 ± 4 | 0.006         |
|                                  | semispinalis capitis | 9              | 33 ± 1              | 7 ± 2                                    | 7 ± 2                                    | 0.001             | 4 ± 2  | 0.033         |

\(a\) \(h^2\) and \(c^2\): coefficients of heritability and of common environment, when it was necessary, used for genetic evaluation by PEST.

\(b\) \(\sigma_p\) and \(\mu_{rn^+}/rn^+\): estimates of the within genotype standard deviation and of the within \(rn^+\)/\(rn^+\) genotype mean (± standard error) computed by SAS GLM.

\(c\) \(\mu_{RN^-}/RN^- - \mu_{rn^+}/rn^+\), \(\mu_{RN^-}/rn^- - \mu_{rn^+}/rn^+\) and \(p1 > \chi^2\): estimates of the contrasts between genotypic means (± standard error) and level of significance of the RN effect (test of the “\(\mu_{RN^-}/RN^- - \mu_{rn^+}/rn^+ = 0\) and \(\mu_{RN^-}/rn^- - \mu_{rn^+}/rn^+ = 0\)” hypothesis) computed by PEST.

\(d\) \(d\) and \(p2 > \chi^2\): estimate (± standard error) \((d = \mu_{RN^-}/rn^- - 0.5 \ (\mu_{RN^-}/RN^- + \mu_{rn^+}/rn^+))\) and level of significance (test of the “\(d = 0\)” hypothesis) of the dominance effect computed by PEST.
3.2.4. Physico-chemical characteristics of muscle

The effects of the RN gene on physico-chemical characteristics of muscle are reported in Table VI. The RN genotype had no effect on *longissimus* pH₁ but, as expected, had a major effect of about 2 s.d. on pH₂₄ of all studied muscles. Furthermore, for the latter trait, the RN⁻ allele appeared to be fully dominant, confirming the non-linear relationship between GP (either I.V. or P.M.) and ultimate pH, with a threshold value of GP beyond which ultimate pH is constant (for review, see [17]).

Effects of RN were also highly significant for most meat colour parameters. The lightness *L*° parameter was increased in RN⁻ carriers by about 0.8 s.d., except for the *biceps femoris* in the RN⁻/rn⁺ genotype, which corresponds to a paler meat. Regarding effects of RN on redness and yellowness, an overdominance situation was encountered, *a*° and *b*° parameters being higher in the heterozygous genotype. Finally, water-holding capacity was decreased in the *longissimus* muscle of the RN⁻ carriers, and there was a similar, but not significant, tendency for the two other muscles studied.

On the whole, preceding hypotheses [7, 22, 23, 30, 33] concerning the implication of the RN gene in the occurrence of the “Hampshire effect” are fully supported by these new observations. On the other hand, the present comparison including homozygous carrier animals allows confirmation of the complete dominance of the RN⁻ allele for most physico-chemical characteristics of meat, e.g. pH₂₄ and colour (*L*° value).

3.2.5. Technological and eating meat quality

As shown in Table VII, most technological and eating meat quality traits were affected by the RN genotype. The estimated difference between homozygotes for Napole yield was 8.4 percentage points (3.2 s.d.), i.e. a value fairly similar to that obtained in the primary study of Le Roy et al. [20]. A difference of around 6 percentage points in Napole yield was reported by Lundström et al. [24] comparing RN⁻/rn⁺ and rn⁻/rn⁺ offspring.

Except for the anatomic yield, the yields measured during the cured-cooked ham processing were lower in the RN⁻ carriers. This decrease was not significant for curing yield, but was highly significant for cooking, technological and overall yields. The difference between homozygotes reached 1.5 s.d. for cooking yield and 1.1 s.d. for technological yield. The RN⁻ allele appeared to be completely dominant over the rn⁺ allele for these traits.

The adverse effect of RN⁻ was of much smaller magnitude for technological yield of cured-cooked ham processing (around 2 percentage points) than for Napole yield (8 percentage points). This is probably related to the peculiar ham processing method used in the present experiment. The process included several phases of tumbling of meat during the curing step resulting in a high average for weight gain at curing (8-9%) and low average values for weight loss during cooking (5-6%). One can assume that such a process led to weakening the differences between RN genotypes. For comparison, Lundström et al. [24], using a similar ham processing method but without tumbling, found an advantage of around 4 percentage points in processing yield for rn⁻/rn⁺ pigs (84.9%), compared with RN⁻/rn⁺ pigs (80.8%).
| Trait      | Muscle                        | $\sigma_p$ | $\mu_{rn^+/rn^-}$ | $\mu_{RN^-/RN^-} - \mu_{rn^+/rn^-}$ | $\mu_{RN^-/rn^-} - \mu_{rn^+/rn^+}$ | $\mu_{RN^-/rn^+} - \mu_{rn^+/rn^-}$ | $p_1 > \chi^2$ | $d^d$           | $p_2 > \chi^2$ |
|-----------|-------------------------------|------------|-------------------|--------------------------------------|--------------------------------------|--------------------------------------|----------------|----------------|----------------|
| pH1       | longissimus                   | 0.17       | 6.59 ± 0.03       | 0.01 ± 0.05                          | 0.02 ± 0.04                          | 0.857                                | 0.01 ± 0.03 | 0.650         |               |
| pH24      | longissimus                   | 0.12       | 5.74 ± 0.02       | -0.20 ± 0.03                         | -0.21 ± 0.03                         | 0.000                                | -0.11 ± 0.02 | 0.000         |               |
|           | gluteus superficialis         | 0.13       | 5.75 ± 0.02       | -0.19 ± 0.03                         | -0.19 ± 0.03                         | 0.000                                | -0.09 ± 0.02 | 0.000         |               |
|           | biceps femoris                | 0.12       | 5.77 ± 0.02       | -0.21 ± 0.03                         | -0.21 ± 0.03                         | 0.000                                | -0.11 ± 0.02 | 0.000         |               |
|           | adductor femoris              | 0.15       | 5.92 ± 0.02       | -0.36 ± 0.04                         | -0.34 ± 0.03                         | 0.000                                | -0.16 ± 0.03 | 0.000         |               |
|           | semispinalis capitis          | 0.13       | 5.75 ± 0.02       | -0.23 ± 0.04                         | -0.21 ± 0.03                         | 0.000                                | -0.09 ± 0.03 | 0.001         |               |
|           | semispinalis capitis          | 0.29       | 6.41 ± 0.05       | -0.31 ± 0.08                         | -0.26 ± 0.08                         | 0.000                                | -0.11 ± 0.06 | 0.091         |               |
| L* (scale 0-100) | longissimus                  | 3.6        | 47.8 ± 0.6        | 2.7 ± 1.0                            | 3.1 ± 0.9                            | 0.002                                | 1.8 ± 0.8    | 0.021         |               |
|           | gluteus superficialis         | 4.2        | 44.3 ± 0.7        | 3.3 ± 1.0                            | 3.3 ± 0.9                            | 0.001                                | 1.7 ± 0.7    | 0.021         |               |
|           | biceps femoris                | 3.6        | 46.5 ± 0.6        | 2.1 ± 0.9                            | 0.8 ± 0.8                            | 0.059                                | -0.3 ± 0.6   | 0.652         |               |
| a*        | longissimus                   | 2.0        | 6.6 ± 0.3         | 1.1 ± 0.5                            | 1.8 ± 0.5                            | 0.002                                | 1.2 ± 0.4    | 0.003         |               |
|           | gluteus superficialis         | 3.6        | 2.4 ± 0.8         | 0.8 ± 0.9                            | 0.5 ± 0.8                            | 0.055                                | 1.5 ± 0.4    | 0.001         |               |
|           | biceps femoris                | 2.6        | 2.3 ± 0.7         | 1.4 ± 0.6                            | 1.0 ± 0.5                            | 0.000                                | 1.0 ± 0.4    | 0.025         |               |
| b*        | longissimus                   | 1.5        | 4.6 ± 0.3         | 0.7 ± 0.5                            | 0.9 ± 0.5                            | 0.384                                | 0.6 ± 0.4    | 0.193         |               |
|           | gluteus superficialis         | 1.6        | 6.6 ± 0.3         | 1.0 ± 0.4                            | 1.6 ± 0.3                            | 0.000                                | 1.1 ± 0.3    | 0.000         |               |
|           | biceps femoris                | 1.8        | 7.7 ± 0.3         | 1.0 ± 0.3                            | 0.9 ± 0.4                            | 0.050                                | 0.6 ± 0.3    | 0.036         |               |
| Water     | longissimus                   | 38         | 182 ± 6           | -28 ± 11                             | -31 ± 10                             | 0.004                                | -17 ± 8      | 0.035         |               |
| holding capacity (s) | longissimus                 | 67         | 122 ± 10          | -15 ± 18                             | -27 ± 16                             | 0.213                                | -20 ± 13     | 0.118         |               |
|           | biceps femoris                | 34         | 184 ± 5           | -6 ± 9                               | -16 ± 7                              | 0.082                                | -13 ± 6      | 0.033         |               |

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$a$ $h^2$ and $c^2$: coefficients of heritability and of common environment, when it was necessary, used for genetic evaluation by PEST.

$b$ $\sigma_p$ and $\mu_{rn^+/rn^-}$: estimates of the within genotype standard deviation and of the within $rn^+/rn^-$ genotype mean (± standard error) computed by SAS GLM.

c $\mu_{RN^-/RN^-} - \mu_{rn^+/rn^-}, \mu_{RN^-/rn^+} - \mu_{rn^+/rn^-}$ and $p_1 > \chi^2$: estimates of the contrasts between genotypic means (± standard error) and level of significance of the RN effect (test of the “$\mu_{RN^-/RN^-} - \mu_{rn^+/rn^-} = 0$ and $\mu_{RN^-/rn^+} - \mu_{rn^+/rn^-} = 0$” hypothesis) computed by PEST.

d $d$ and $p_2 > \chi^2$: estimate (± standard error) ($d = \mu_{RN^-/rn^+} - 0.5 (\mu_{RN^-/RN^-} + \mu_{rn^+/rn^-})$) and level of significance (test of the “$d = 0$” hypothesis) of the dominance effect computed by PEST.
Table VII. Effect of the RN genotype on technological and eating meat qualitya.

| Trait                              | $\sigma_p$ | $\mu_{m^+/m^+}$ | $\mu_{RN^-/RN^-} - \mu_{m^+/m^+}$ | $\mu_{RN^-/m^+} - \mu_{m^+/m^+}$ | $p_1 > \chi^2$ | $d$ | $p_2 > \chi^2$ |
|------------------------------------|------------|-----------------|-----------------------------------|----------------------------------|----------------|-----|--------------|
| Cured-cooked ham processing ability|            |                 |                                   |                                  |                |     |              |
| Anatomic yield (%)                 | 2.5        | 74.3 ± 0.4      | 0.1 ± 0.8                         | -0.3 ± 0.7                       | 0.826          | -0.3 ± 0.5 | 0.541        |
| Curing yield (%)                   | 1.7        | 109.0 ± 0.3     | -0.6 ± 0.5                        | -0.8 ± 0.5                       | 0.194          | -0.5 ± 0.4 | 0.174        |
| Cooking yield (%)                  | 1.0        | 95.2 ± 0.2      | -1.5 ± 0.3                        | -1.3 ± 0.3                       | 0.000          | -0.5 ± 0.2 | 0.011        |
| Technological yield (%)            | 2.0        | 103.8 ± 0.3     | -2.2 ± 0.6                        | -2.2 ± 0.5                       | 0.000          | -1.1 ± 0.4 | 0.012        |
| Overall yield (%)                  | 2.6        | 77.1 ± 0.4      | -1.6 ± 0.8                        | -1.9 ± 0.7                       | 0.017          | -1.1 ± 0.6 | 0.045        |
| Eating quality of grilled chops    |            |                 |                                   |                                  |                |     |              |
| Visual compactness at cutting (0-10)| 1.8        | 5.5 ± 0.1       | 0.1 ± 0.4                         | 0.8 ± 0.4                        | 0.059          | 0.7 ± 0.3 | 0.018        |
| Tenderness (0-10)                  | 2.0        | 6.3 ± 0.2       | -0.9 ± 0.5                        | -1.2 ± 0.4                       | 0.015          | -0.7 ± 0.4 | 0.035        |
| Juiciness (0-10)                   | 1.9        | 5.8 ± 0.2       | -0.2 ± 0.4                        | -0.6 ± 0.4                       | 0.268          | -0.5 ± 0.3 | 0.122        |
| Mellowness (0-10)                  | 1.9        | 5.7 ± 0.2       | -0.8 ± 0.5                        | -0.8 ± 0.4                       | 0.096          | -0.5 ± 0.3 | 0.174        |
| Pork flavour intensity (0-10)      | 1.8        | 4.7 ± 0.1       | 1.4 ± 0.4                         | 0.9 ± 0.4                        | 0.003          | 0.2 ± 0.3 | 0.620        |

a Coefficients of heritability and of common environment, when it was necessary, used for genetic evaluation by PEST: $h^2 = 0.30$ and $c^2 = 0.05$ for Napole yield; $h^2 = 0.50$ for anatomic yield and $h^2 = 0.40$ for curing, cooking, technological and overall yields; $h2 = 0.20$ for eating meat quality descriptors.

b $\sigma_p$ and $\mu_{m^+/m^+}$: estimates of the within genotype standard deviation and of the within $m^+/m^+$ genotype mean (± standard error) computed by SAS GLM.

c $\mu_{RN^-/RN^-} - \mu_{m^+/m^+}$, $\mu_{RN^-/m^+} - \mu_{m^+/m^+}$ and $p_1 > \chi^2$: estimates of the contrasts between genotypic means (± standard error) and level of significance of the RN effect (test of the $\mu_{RN^-/RN^-} - \mu_{m^+/m^+} = 0$ and $\mu_{RN^-/m^+} - \mu_{m^+/m^+} = 0$ hypothesis) computed by PEST.

d $d$ and $p_2 > \chi^2$: estimate (± standard error) ($d = \mu_{RN^-/m^+} - 0.5 (\mu_{RN^-/RN^-} + \mu_{m^+/m^+})$) and level of significance (test of the $d = 0$ hypothesis) of the dominance effect computed by PEST.
Some of the eating quality traits were also influenced by the RN gene. Score for tenderness was lower in the RN− carriers while these animals exhibited pork flavour intensity. The RN−/RN− and RN−/rn+ were close to each other for tenderness. The gene effect on flavour was approximately additive. Lundström et al. [23] found also a superiority of RN− carriers for meat taste and smell intensities. However, opposite to our results, no difference between RN−/rn+ and rn+/rn+ animals was found for tenderness in Swedish studies and a slight decrease of shear force [23] or chewing time [24] was even shown by RN− carriers. This discrepancy could be explained by difference in the variability of QTL linked to the RN gene between the two experiments: the observed effect on tenderness of meat is perhaps not a pleiotropic effect of the RN gene but an effect of one or several loci closely linked to RN.

4. CONCLUSION

Numerical comparison between estimates, given either by full pedigree analysis or by simpler analysis of the diallel design considering offspring genotypes as fixed effects, supported the hypothesis that the parents used in diallel matings were correctly genotyped and sampled, as regards their polygenic value. A fortiori, estimates of RN genotype effect on RTN-correlated (or non-correlated) traits may be considered as only slightly biased (less than 5%) by miss-genotyping or selection influence. In the present article, RN effects have thus been evaluated from the performance records of the diallel offspring only, using “classical” animal model procedures.

This study definitely confirms that the porcine RN gene has considerable effects on muscle glycolytic potential and some GP-related traits. When expressed in standard deviation unit of the trait, the effect of RN on GP of longissimus muscle is found to be comparable in magnitude with the largest single-gene effects currently known in animals, e.g. dwarfing genes in mouse and chickens, the muscle hypertrophy gene in cattle and the “Booroola” gene in sheep. The identity between the acid meat characteristics and the RN− allele effect is clearly demonstrated. The main features of the “Hampshire” effect were observed in RN− carrier animals: higher muscle GP, lower ultimate pH, paler meat and lower protein content. Moreover, our results are fully consistent with those previously obtained from comparing “high GP” and “low GP” Hampshire-cross pigs even though one discrepancy was noted for tenderness of meat. Furthermore the present comparison between the three RN genotypes allowed to confirm the complete dominance of the RN− allele for RTN, and most meat quality traits. However, the dominance was not quite complete for IVGP which is probably the primary trait affected by the RN gene. Finally, the relationship between GP and carcass lean to fat ratio was confirmed here in the frame of the RN gene segregation. This association could arise either from a pleiotropic effect of the RN gene itself or from effects of other loci located close to RN and in linkage disequilibrium with RN in the present population of sires.
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