Review

Candidate Genes in Bull Semen Production Traits: An Information Approach Review

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Abstract: Semen quality plays a crucial role in the successful implementation of breeding programs, especially where artificial insemination (AI) is practiced. Bulls with good semen traits have good fertility and can produce a volume of semen per ejaculation. The aim of this review is to use an information approach to highlight candidate genes and their relation to bull semen production traits. The use of genome-wide association studies (GWAS) has been demonstrated to be successful in identifying genomic regions and individual variations associated with production traits. Studies have reported over 40 genes associated with semen traits using Illumina BeadChip single-nucleotide polymorphism (SNPs).

Keywords: semen production parameters; candidate genes; genome-wide association studies; bull

1. Introduction

Semen traits are important for cattle gene improvement programs relying on artificial insemination [1]. The collection of bull semen and the scoring of parameters has become the standard routine in many AI centers [2]. Semen production traits that are routinely assessed from fresh ejaculates include semen volume (VOL), sperm concentration (SCONC), sperm motility (SM), and the proportion of sperm with a head (PSH). These traits have key value in assessing semen in AI companies and are also essential when selecting the top-performing bulls [3]. Young bulls are used in AI programs for a short time (4–6 months), which again highlights the importance of semen production traits [3].

Studies [4,5] have focused on identifying genes and genetic markers associated with bovine semen traits in order to understand their genetic architectures. A number of studies on candidate genes associated with semen quality traits have been conducted in pigs [6,7], horses [8], sheep [9], and goats [10,11]. However, the population sizes analyzed in these studies were not large (139–900), with low statistical power to detect causal genes. A small number of genes are shared between these studies; therefore, the genetic structure underlying semen traits remains unknown. Genome-wide association studies can help to overcome the limitations by using single-nucleotide polymorphism (SNP) genotyping assay tools to examine traits of significance [12]. Fortes et al. (2013) [13] revealed that significant SNPs in the X-chromosome were associated with the percentage of progressive motile spermatozoa at 18 months of age and the percentage of normal spermatozoa at 24 months of age in tropical composite bulls. A study conducted by Gottschalk et al. (2016) [8] on 139 German Warmblood horses identified that 29 SNPs on 12 unique chromosomes
were associated with semen quality traits. A number of studies have identified candidate genes in livestock and continue to demonstrate the efficiency of GWASs to identify semen traits [14–16]. Hering et al. (2014) [14] conducted GWASs on 41 bulls with very poor sperm motility and 279 bulls with excellent sperm motility and identified nine candidate genes all with a strong relationship to sperm function. Hence, this review uses the information approach to highlight candidate genes, and their relation to bull semen production traits.

2. Association Studies for Semen Traits

A genome-wide association study (GWAS) is an approach used in genomic research to associate specific genotype variations with a particular phenotypic trait. Genome-wide association studies evaluate genomes from several phenotypes, looking for genetic markers that can be used to predict the presence of a trait. Once such markers are identified, they can be used to understand how genes contribute to the traits. Genome-wide association studies have several factors that contribute to their successful utility. One example is the phenotypic variation; filtering phenotype data is important to avoid outliers during analysis. The population size is very important for obtaining meaningful results for both phenotypic and genotypic variation; one example is a recent study on 1819 Angus bulls with 50,624 records for a single-step genome-wide association study (ssGWAS) on the following traits: VOL, SCONC, number of spermatozoa (NS), initial motility (IMOT), post-thaw motility (PTMOT), three-hour post-thaw motility (3HRPTMot), percentage of normal spermatozoa (%NORM), primary abnormalities (PRIM), and secondary abnormalities (SECs) [16]. Their findings indicated regions of the genome that impacted fertility and included genomic information in the genetic evaluation, which is advantageous for genetically improving male fertility traits [16]. Single-nucleotide polymorphisms (SNPs) are bi-allelic genetic markers, easy to evaluate and interpret, and widely distributed within genomes. These SNPs can compute linkage disequilibrium (LD), fixed information in the genome used to classify fundamental genes of adaptation in domestic animals. Genome-wide studies have been conducted in both beef and dairy cattle and have revealed several genomic regions associated with semen traits.

Hering et al. (2014) [14] reported markers in Holstein Friesian bulls that were significant for semen volumes on BTA 10 and 22: rs42438348, rs41625599, rs41584616, and rs42012507 and the total number of motilities on BTA 22 rs41625599, rs41584616, rs42012507, and rs110109069. They further indicated that only 19 SNPs were significantly associated with five semen traits. Recently, studies of association have depended on the genotyping of thousands of SNP markers to evaluate genomic breeding value. One study mapped 34 SNPs in 19 different Bos taurus autosomes and one on chromosome X: significant SNPs were located on chromosome 24 (rs110876480), 5 (rs110827324 and rs29011704), and 1 (rs110596818), in close vicinity to the melanocortin 4 receptor (MC4R) gene, the PDZ domain containing the ring finger 4 (PDZRN4) gene, the ethanolamine kinase 1 (ETNK1) gene, and the olfactory receptor 5K3-like (LOC785875) gene, respectively [15]. The cost of processing GWASs is reasonable, due to the low cost of sequencing and the ability to determine large datasets. GWASs require a bi-allelic assumption, which is reasonable because only 1–3% of the genome has random copy number variants. Liu et al. (2017) [17] reported that one SNP (rs110305039) located on downstream of PDGFRB was significantly related to semen volume per ejaculate (SVPE), the number of sperm per ejaculate (NSPE), and the number of motile sperm per ejaculate (NMSPE). This SNP rs110305039 was previously revealed to have an association with sperm motility (SM). Liu et al. (2017) [17] further revealed three significant SNPs, rs211260176, rs208093284, and rs43445726, located on the promoter of MARCH1, which showed associations with the semen volume per ejaculate (SVPE), number of sperm per ejaculate (NSPE), and number of motile sperm per ejaculate (NMSPE), respectively. Figure 1 shows the distribution of the 1364258 bp length start and 19755465 bp length end SNP on the chromosome, which was taken from our GWAS data using QTL/association. Recently, Butler et al. (2022) [18] reported five SNPs, BTB-01549373, rs4166488, rs109736826, rs109268478, and rs41575945, which are strongly associated with
SV, and three significant SNPs, rs43067163, rs41623602, and rs29023737, which were identified for influencing concentration (CONC) [18]. Moreover, for IMot, six SNPs on six different chromosomes were significant, including rs41623436, rs109798673, rs43526428, rs29003479, rs42861585, and rs109512383 [18]. For % NORM, six significant SNP were identified: rs41606310 on chromosome one, rs110964837 on chromosome two, rs41594758 on chromosome three rs410928164 on chromosome five, rs41591913 on chromosome five, and rs41666416 on chromosome ten [18].

![Figure 1](https://www.animalgenome.org/cgi-bin/QTLdb/BT/search accessed on 29 January 2022)

**Figure 1.** Length of marker position of a gene (MARCH1) and Chromosome 6 (https://www.animalgenome.org/cgi-bin/QTLdb/BT/search accessed on 29 January 2022).

### 2.1. Single-Nucleotide Polymorphism Markers Used to Identify Associations

Genome-wide association studies have relied on certain types of single-nucleotide polymorphism markers to obtain information in different populations and species. The SNPs selected in GWASs are either Illumina (San Diego, CA, USA) or Affymetrix (Santa Clara, CA, USA). These two contending technologies have been selected and reviewed and bring unique approaches to measure SNP differences. Affymetrix (chip-based) displays short DNA sequences to detect alleles, whereas Illumina (bead-based) displays slightly longer DNA sequences to detect alleles. A study by Suchocki and Szyda (2015) [4] reported that, recently, most genome-wide association studies on fertility traits have been conducted using the bovine 50K Illumina SNP chip. The information gathered by these markers can assist in identifying phenotypic differences and highlighting the accuracy of animal selection for mating purposes, thus increasing genetic gains for advanced selection pressures [19]. However, numerous association studies conducted on complex traits have used Illumina BeadChip SNPs, especially in studies identifying semen production traits. A summary of studies conducted using Illumina BeadChip SNPs is presented in Table 1.

### Table 1. Studies conducted using Illumina on semen traits.

| Population Size          | Target Traits                                                                            | SNPs Used                                  | Articles Published |
|--------------------------|------------------------------------------------------------------------------------------|--------------------------------------------|--------------------|
| 1581 Holstein Friesian   | Sperm motility                                                                           | Illumina BovineSNP50K BeadChip, 54,001.   | [3]                |
| 1212 bulls               | Sperm concentration, semen volume, number of spermatozoa, motility, and motility score.  | Illumina BovineSNP50K BeadChip, 54,001.    | [4]                |
| 692 Holstein bulls       | Ejaculate volume, sperm concentration, sperm motility, and sperm concentration            | Illumina BovineSNP50 BeadChip, SNP markers | [5]                |
| 1085 tropical composite  | Sperm motility and percentage of normal spermatozoa.                                     | Illumina Bovine HD SNP50 BeadChip          | [13]               |
| 788, Holstein Friesian   | Semen volume and total number of sperm                                                   | Illumina BovineSNP50 Bead-Chip, 54,001.    | [14]               |
### Table 1. Cont.

| Population Size | Target Traits                                                                 | Snps Used                              | Articles Published |
|-----------------|-------------------------------------------------------------------------------|----------------------------------------|-------------------|
| 730 Chinese Holstein bulls | Semen volume per ejaculate, sperm motility, sperm concentration per ejaculate, the number of sperms per ejaculate, and the number of motile sperms per ejaculate. | Illumina Bovine SNP50 BeadChip          | [17]              |
| 1819 Angus      | Volume, concentration, number of spermatozoa, initial motility, post-thaw motility, three-hour post-thaw motility, percentage of normal spermatozoa, primary abnormalities, and secondary abnormalities. | Illumina Bovine HD SNP50 BeadChip      | [18]              |
| 1799, Austrian Fleckvieh bulls | Total number of spermatozoa and percentage of live spermatozoa. Sperm morphology and sperm chromatin phenotypes. Ejaculate volume, sperm concentration, sperm motility, sperm head, and tail anomalies. | Illumina Bovine SNP50K BeadChip 54,001. Illumina Bovine SNP50 BeadChip 54,000. | [20] [21]        |
| 1099 Brahman and Tropical bulls 1719 |                                                      |                                        |                   |
| 2481 Brown Swiss bulls |                                                      |                                        |                   |

#### 2.2. Candidate Genes Associated with Semen Traits

The candidate gene approach is useful for quickly determining the associations of a specific genetic variant with a phenotype and the number of relevant genes governing traits [23]. To date, several candidate genes have been detected that influence valuable traits, even though the total amount of published information of putative genes remains quite small. The candidate genes approach has been criticized due to the low reputability of results and the limitations of including relevant genes [24]. The challenge is that it requires existing information on physiological, biochemical, or functional knowledge and the biochemical metabolism pathway, which may not be available [25]. Candidate gene association studies evaluate genetic variations associated with traits within a limited number of pre-specified genes [26]. For instance, candidate gene studies have more significant outputs in identifying variation, critical genes, and biological pathways [26].

Many putative candidate genes in both dairy and beef cattle have been identified to be associated with bull semen traits. Studies on genes of biological pathways showed an associated with semen traits, which revealed seven significant biological pathways relating to 127 genes with 1.04% of the genetic variation for the volume (VOL), number of sperm (NS), and motility (MOT) [27]. Nonetheless, studies on candidate genes have been performed, and several genetic variations have been identified in bull semen. Borowska et al. (2018) [28] reported that candidate genes had two shared SNPs related to sperm plasma integrity and used GWASs for SC records on 392 bulls and identified the STATU2 gene which participates in multiple biological processes, including reproduction and developmental and immune structures. Hering, Olenski, and Kaminski (2014) [3] used two approaches to find candidate genes potentially related to very poor sperm motility: (1) a physical approach, which involved genes in the vicinity of a significant pathway, an approach that identified nine candidate genes positioned close to significant SNP markers found to have a strong relationship with sperm function (LOC785875, ALPL, HIBADH, GADD45G, TRIM36, TGFA, KLHL1, PRKCB, and INCENP); (2) a functional approach, which is the identification of candidate genes with a close distance of 1 mb. These strategies can assist in identifying candidate genes associated with important traits of interest. The genes identified above support the hypothesis that looking for candidate genes only among those in the direct vicinity of significant markers can overlook other genes. Table 2 shows the genes found in several studies of BTA positions and SNPs of interest. Genes such as FSHR, INHA, TNP1, TNP2, CAPN1, and SPAG11 have been studied as candidate genes...
for influencing semen traits in bulls [25, 29], as shown in Tables 2 and 3. The benefits of sequencing a whole bovine genome [25] with many SNPs can simplify the localization of specific gene regions associated with traits of importance. A study on genomic regions related to the inbreeding depression of live spermatozoa identified 53 genes, and an additional analysis of efficient annotations of the genes recognized nine strong candidate genes related to male fertility, located on chromosomes 1, 6, 10, and 14 [30]. However, there appears to have been surprisingly little work conducted to date to characterize bovine Y chromosome genes for male fertility [27]. This is probably because, until recently, there has been no reference sequence assembly for the bovine Y chromosome [27]. One candidate gene commonly found to be associated with bull sperm quality is the SOX5 (SRY-box5) gene, which encodes for the sex-determining Y chromosome box5 protein [28]. Five of the prioritized candidate genes for sperm motility have also been formerly revealed as candidate genes in the regulation of male fertility. These include superoxide dismutase 2, T-complex protein 1, parkin co-regulated gene, sperm flagella 2 gene, and prolactin receptor (SOD2, TCP1, ACRG, SPEF2, and PRLR). One study reported on 22 novel candidate genes which were identified on chromosomes 1, 5, 6, 7, 15, 17, 23, and 27 [5]. There are many more window regions affecting traits, which explain up to 1% of genetic variance. Results from Qin et al. (2017) [5] exhibited candidate genes PDE3A and SLCO1C1, where PDE3A was on BTA5. Candidate gene studies can overcome these issues, focusing directly on the association between disease and variants in particular genes that have a priori biological support. This focus comes at a cost: candidate gene studies ignore much of the genome and thus are likely to miss many causal regions or genes and instead find many false-positive associations.

Table 2. Candidate genes, markers, and chromosome positions [3].

| SNP Name       | Position | Identified Candidate Genes | Reference |
|----------------|----------|----------------------------|-----------|
| rs29010277     | X        | LOC100848828               | [4]       |
| rs110419531    | X        | MAGEB10                    | [4]       |
| rs109349108    | X        | MAGEB10                    | [4]       |
| rs110685046    | X        | MAGEB10                    | [4]       |
| rs211260176    | 6        | MARCH1                     | [17]      |
| rs211260176    | 6        | MARCH1                     | [17]      |
| rs110128350    | 27       | DLC1                       | [23]      |
| rs109170505    | 22       | OPN1LW                     | [31]      |
| rs110876480    | 1        | GABRR3                     | [20]      |
| rs109466217    | 2        | EF4G3                      | [32]      |
| rs109416157    | 29       | INCENP                     | [33]      |
| rs43399120     | 4        | CFTR                       | [25]      |
| rs109697710    | 4        | HIBADH                     | [34]      |
| rs290117704    | 5        | SOX5                       | [35]      |
| rs42601646     | 5        | PTPRB                      | [36]      |
| rs2601646      | 5        | PTPRR                      | [37]      |
| rs11065449     | 8        | SECISBP2                   | [37]      |
| rs42749302     | 8        | CYCL2                      | [36]      |
| rs109677705    | 11       | SPAST                      | [38]      |
Table 2. Cont.

| SNP Name   | Position | Identified Candidate Genes | Reference |
|------------|----------|----------------------------|-----------|
| rs41574912 | 12       | KLHL1                      | [39]      |
| rs41965546 | 21       | TGFIR                      | [40]      |
| rs110876480| 24       | GRP                        | [41]      |
| rs110149073| 25       | PRKCB                      | [42]      |
| rs42736384 | 26       | HELLS                      | [42]      |
| rs109339115| 29       | CATSPERR1                  | [24]      |
| rs109339115| 29       | CAPN1                      | [43]      |

Table 3. Genes associated with different semen parameters.

| Associated Trait                  | Chromosomes | Identified Candidate Genes | Reference |
|-----------------------------------|-------------|---------------------------|-----------|
| semen volume per ejaculate        | 6           | MARCH1                    | [17]      |
| number of motile sperm per ejaculate | 6           | MARCH1                    | [17]      |
| Sperm motility                    | 6           | MARCH1                    | [17]      |
| Percentage of live sperm          | 1           | SPATA16                   | [31]      |
| Total number of sperms            | 14          | RPL10L                    | [31]      |
|                                   | 1           | NYD-SP5                   | [31]      |
|                                   | 10          | SPESP1                    | [31]      |
| Semen motility                    | 27          | COX7A2L                   | [31]      |
|                                   | 25          | DNAH3                     | [44]      |
|                                   | 5           | PRP11                     | [45]      |
| Ejaculate volume                  | 10          | PSMB5                     | [46]      |
|                                   | 16          | NR5A2                     | [46]      |
|                                   | 10          | PRMT5                     | [46]      |
| Sperm concentration               | 25          | FSCN1                     | [46]      |
| Number of motile sperm            | 24          | IQCJ                      | [46]      |
| Number of sperms per ejaculate    | 3           | LHX8                      | [46]      |
|                                   | 24          | NPC1                      | [46]      |
|                                   | 8           | DMRT1                     | [46]      |
| EV, SPC, TSN, SM, and PTM         | Y           | ZNF280AY                  | [47]      |
| Semen quality                     | Y           | SOX5 (SRY-box5)           | [28]      |

2.3. Genes Detected in Dairy and Beef Cattle

The identification of candidate genes provides a better understanding of the distribution of genes that affect traits of economic interest [48]. Hering et al. (2014) [3] reported on the role of the following genes associated with sperm motility using a physical approach in a Holstein population: MC4R, ETNK1, LOC785875, TRIM36, ALPL, PRKCB, HIBADH, KHL1, PD2RN4, CTR, SRD5A2, CAPN1, CATSPER1, ATP5O, GABRR3, EIF4G3, SOX5, PTPRB, SECISBP2, CYLC2, CRYZL1, LOC785875, ZBTB40, ALPL, AGBL4, ST7, PDZRN4, CNOT2, ETNK1, MRPL1, RAPGEF6, FREM1, GADD45G, SPIN1, GRIN3A, TRIM36, LOC511898, TGFA, FAM84B, LOC10139627, CYP2C87, SORCS1, LOC101905219, and DLC1 (shown in Table 4). In another study, Hering et al. (2014) [14] highlighted genes DCP1, SFMBT1, GALC, PRKCD, PHF7, TLR9, SPATA7, and TMEM110 associated with semen volume and the total number of sperm in Holstein Friesian bulls, located in the vicinity of significant markers. However, most studies on candidate gene association have strongly focused on dairy cattle, as shown in Table 4. For beef cattle, however, there is limited information on candidate gene association studies. Sweett et al. (2020) [16] identified five candidate genes in 265 crossbred beef bulls for SM in the regulation of male fertility. These genes include superoxide dismutase 2, T-complex protein 1, parkin co-regulated gene, sperm flagella 2 gene, and prolactin receptor (SOD2, TCP1, PACRG, SPEF2, and PRLR). Other results on beef cattle have been documented by Butler et al. (2022) [18]; genes...
associated with fertility were found to be close to the significant SNPs in the study. There is still a gap where dairy cattle research has capitalized on genomic technologies [3,49] and multiple QTL regions [5,46] and candidate genes [50,51] associated with male and female fertility have been identified. There is limited information on fertility traits in beef cattle bulls [16]. This gap was highlighted by studies conducted by Sweett et al. (2020) [16] and Butler et al. (2022) [18], which showed a lack of attention given to beef cattle regarding candidate genes. Sweett et al. (2020) [16] even stated that SM measurements are much less common in the beef industry, where natural breeding is often used, as can be seen in our sample size.

Table 4. Breeds, genes, and traits of association.

| Breeds                        | Genes                                           | Traits            | Reference |
|-------------------------------|-------------------------------------------------|-------------------|-----------|
| Dairy breed, chinese holstein  | ETNK1, PDE3A, CSF1R, WTI, DSCAML1, SOD1, and RUNX| Semen traits      | [5]       |
|                               | MARCH1, PDGFRB, and PDE3A                       | SVPE, SCPE, NSPE, and NMSPE | [17]      |
|                               | PSMS, PRMT5, ACTB, PBDE3A, FSCN1, NR5A2, IQCG, LANX8, and DMRT1 | VE, SM, SC, NSP, and NMSP | [46]      |
|                               | M4C4R, ETNK1, LOC785875, TRIM36, ALPL, PRKCB, HIBADH, KHL1, PD2RN4, CTRR, SRD5A2, CAPN1, CATSPE1, ATP5O, GABBR3 EIF4G3, SOX5 PTPRB PTPRR, SECSB2, CYLC2, CRYZL1, LOC785875, ZBTB40, ALPL, AGB14 ST7, PD2RN4, CNOT2, ETNK1 MRPL1, RAPGEF6, FREM1 GADD45G, SPIN1 GRIN3A TRIM36 LOC511898 TGFA, FAM84B, LOC100139627, CYP2C87, SORCS1, LOC101905219, and DLC1 | Poor SM | [3,32–47] |
|                               | FSHR, INHBA, INHA, and PRL                       | VOL, SCON, MOT, FMOT, AIR, and ASR | [29]     |
| Brown swiss                   | WDR19                                           | Semen quality     | [22]     |
| Dual purpose breeds,          | DCP1, SFMBT1, GALC, PRKCD, PHF7, TLR9, SPATA7, and TMEM110 | SV and TNP | [14] |
| holstein friesian             | PDRK2 and GALNT113                              |                   |           |
| Beef breeds, crossbreeds      | MAGEB10 and KLHL113                             | SV, MS, M, SC, and NS | [4] |
| (angus, simmenthal,           | WTAP, ACAT2, TCP1, EZR, PRKN, PRC3G, PLCB4, LAMP5, PAK5LMBRD2, UGT3A2, CAPSL, IL7R, SPEF2, SKP2, PRLR, and DCC | SM | [16] |
| piedmontese, gelbvieh,        |                                                  |                   |           |
| charolais, and limousine.)     |                                                  |                   |           |
| American angus                | HERC2, OCA2, and LOC101902976                   | VOL, CONC, NSP, IMot, P3RTM3, 3HRPTMot, %NORM, PRIM, and SEC | [18] |

Some of these genes were reported to have a significant role in spermatogenesis, as shown in Table 4. The genes LOC785875, ALPL, HIBADH, GADD45G, TRIM36, TGFA, KLHL1, PRKCB, and INCENP have a strong relationship with sperm function, which confirms the important role of these genes on semen quality [3]. Table 5 represent all the genes found in GWAS studies and in my review paper.
3. Traits in Most GWAS Studies

Semen traits are easily measurable, and record-keeping for each bull is essential in any AI company. Most of the studies conducted on bull semen traits have focused on several important parameters, such as motility, progressive motility, and morphological abnormalities of sperm [3,17,62]. These are some of the parameters regarded as critical in assessing semen by AI companies. However, selecting animals directly based on their semen phenotypes can be difficult because of the low (0.04) to moderate (0.30) heritability of these traits [49,63]. Gredler et al. (2007) [63] demonstrated the moderate heritability of all semen parameters with a low correlation between breeding values for semen quality traits, and routinely estimated the breeding values for male fertility. Stålhammar et al. (1989) [64] highlighted the low to moderate heritability of semen parameters in Swedish Red, White, and Swedish Friesian bulls. The demand for bulls with good semen parameters is very high; hence, regular production should be a priority for AI companies [30]. Studies in the literature have shown moderate to high heritability for some of the parameters, whereas in other studies, heritability is lower, making it hard to select the same parameters in other animals. Yin et al. (2019) [46] studied the heritability of the following semen parameters: ejaculate volume (VE), progressive sperm motility (SM), sperm concentration (SC), number of sperm (NSP), and number of motile sperm per ejaculation and the number of motile sperm per ejaculate. However, there is limited information on some of the parameters in these studies; most of these evaluated parameters are repeated across studies. Karoui et al. (2011) [65] revealed moderate heritability for the volume, concentration, number of spermatozoa per ejaculate, mass motility score, and post-thawing motility traits; only individual motility was low in heritability. A more inclusive view regarding the semen traits to be measured across all parameters could broaden bull selection criteria in the future. This can expand knowledge of the importance of semen production parameters in bulls. Additionally, to improve the power of the analysis, multi-trait techniques and denser marker maps are required for reference for future studies.

Table 5. Genes identified in this GWAS studies and their full names.

| Gene ID   | Gene Full Name                                  | Gene ID   | Gene Full Name                  |
|-----------|-----------------------------------------------|-----------|---------------------------------|
| TGFA      | Transforming growth factor, Ralph [3]          | PRKCB     | Protein kinase C, beta [3,23]    |
| ETNK1     | Ethanolamine kinase 1 [1,5]                    | HIBADH    | 3-Hydroxyisobutyrate dehydrogenase [3,33] |
| LOC785875 | Olfactory receptor SK3-like [3,53]             | INCENP    | Inner centromere protein antigens [3,136] |
| ALPL      | Alkaline phosphatase, liver/bone/kidney [54]  | SOD1      | Superoxide dismutase 1 [5]      |
| TRIM36    | Tripartite motif containing 36 [16]            | ACAT2     | Acetyl-CoA Acetyltransferase 2 [16] |
| MC4R      | Melanocortin 4 receptor [15]                   | TCP1      | T-complex protein 1 subunit alpha [16] |
| RSPH3     | Radial Spoke Head 3 [16]                       | GABRR3    | Gamma-Aminobutyric Acid Type A Receptor Subunit Rh3 [3,23] |
| TAGAP     | T Cell Activation RhoGTPase Activating Protein [16] | CAPN1    | Calpain 1 (mu/l) large subunit [43] |
| FNDC1     | Fibrinectin Type II Domain Containing [16]     | CATSPER1  | Cation channel, sperm-associated 1 [3,23] |
| F9        | Coagulation factor IX [55]                     | ATGPSO    | ATP synthase, H+ transporting, mitochondrial [3,5,34] |
| WDRED1    | WD Repeat Domain 19 [22]                      | EIF4G3    | Eukaryotic translation initiation factor 4 gamma, 3 [3] |
| PTPRR     | Protein tyrosine phosphatase, receptor type, R [3,36] | CYCL2    | Cyclic, basic protein of sperm head cytoskeleton 2 [3,56] |
| SOR5      | SRK-Box Transcription Factor 5 [5,35]          | CRYZ1L    | CRY2-like [3] |
| PTPR8     | Protein tyrosine phosphatase, receptor type, B [3,36] | LOC785875 | SECIS binding protein 2Cylicin, basic protein of sperm head cytoskeleton 2 [57] |
| SECISBP2  | secIS binding protein 2Cylicin, basic protein of sperm head cytoskeleton 2 [57] | OCA2     | oculocutaneous albinism 1 [57] |
| KLHL1     | kelch-like 1 (Drosophila) [59]                 | AGBL4     | ATP/GTP binding protein-like 4 [3] |
| PDGFRB    | Platelet-derived growth factor receptor beta [5,17,58] | RAPGFP6  | Rap quinine nucleotide exchange factor (GEF) 6 [3] |
| PDE3A     | phosphodiesterase 3A [7,28]                   | ZBTB40    | Zinc finger and BTB domain containing 40 [3] |
| DCPI      | decapping mRNA 1A [16]                        | ST7       | Suppression of tumorgenicity 7 [3] |
| SFMBT1    | Scm-like with four mbt domains 1 [14]          | CNOT2     | CC44-NOT transcription complex, subunit 2 [3] |
| PRKCD     | protein kinase C [14]                         | ETRNK C   | Mitochondrial ribosomal protein L1 [3] |
| KLH13     | Kelch-Like Family Member 13 [4]               | MRPL1     |                  |
| HERC2     | HECT and RLD Domain Containing E3 Ubiquitin    | FREM1     | FRAS1 related extracellular matrix [3] |
| MARCH1    | membrane-associated ring finger 1 [17,59]     | SPIN1     | spindlin 1 [3] |
| PYH7      | PH finger protein 7 [40]                      | KIBRE3    | Kirre-like nephrin family adhesion molecule 3 [56] |
| GALK      | Galactosyltransferase [61]                    | LOC51898  | Protein disulfide isomerase family A, member 6 [3] |
| TLR9      | Toll-like receptor 9 [56]                     | SPATA7    | spermato genesis associated 7 [3] |
| TMEM110   | transmembrane protein 110 [14]                | NRS2A     | Nuclear Receptor Subfamily 5 Group A [46] |
|            |                                               | MAGEB10   | melanoma antigen family B10 [4] |

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4. Conclusions

Our review has highlighted genes associated with semen parameters; over 40 genes have been documented in the literature and have been reported by several authors to be associated with semen traits. We have furthered the understanding of how the length of the marker highlights the position of a gene to influence a trait in chromosomes. Marker efficiency is vital in candidate association studies in which Illumina is the most widely used marker.

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