## Effect of genomic pre-selection on the stability of EBVs from traditional BLUP procedure for production traits - a practical illustration

U S Nielsen<sup>1</sup>, J Pösö<sup>2</sup>, P Madsen<sup>3</sup>, E A Mäntysaari<sup>4</sup>, J Pedersen<sup>1</sup>, G Su<sup>3</sup>, G P Aamand<sup>5</sup>

> <sup>1</sup>Seges, Cattle, Denmark <sup>2</sup> Faba co-op, Finland

<sup>3</sup> Centre for Quantitative Genetics and Genomics, Aarhus University, Denmark <sup>4</sup>Luke, Finland, <sup>5</sup> Nordic Cattle Genetic Evaluation, Denmark





# Current status for genomic selection HOL, (DNK,FIN,SWE)

- One out of ten bulls selected for total merit (selection intensity 1.8 and selection differential 1.4)
- Selection intensity for the single traits in total merit is lower (Protein: selection intensity 1.0 selection differential 0.8)
- Second batch bulls are used less intensively. Only 10-20% of inseminations is done with second batch bulls.
- Genomic selected bulls are used intensively as bull sires
- Progeny test by traditional BLUP procedure is done per trait or trait group





## Traditional BLUP evaluation

Unbiased, if all data on which selection is based is included

Ignoring genomic selection

#### In theory:

- Evaluations are biased because in BLUP,, progenies are assumed to be average of their parents, i.e. E[MS-term]=0
- Leads into problems of separating environmental and genetic levels, i.e. genetic trend is underestimated and environmental trend is overestimated
- as a result: young bulls in progeny test are underestimated



# Question: how robust is our evaluation model against unaccounted selection?

- Phenotypic trend = Environmental + genetic trends
- Genetic trend estimation in BLUP:
  - Selection of parents (and now Mendelian term)
- Environmental trend estimation in BLUP:
  - Animals from different generations/birth years are producing in same environment classes
  - 1. First crop vs. second crop daughters
  - 2. Same bulls having daughters in consecutive years
  - 3. Same cows having records in consecutive years

## Objective

- Test the effect of bull second crop daughters for the robustness
- Try to test the effect of genomic selection on accuracy of evaluations
- Mimic the effect of bias from pre-selection on real data for a strongly selected trait. Protein is chosen





## Material

- Nordic Test Day model for production trait (Multi trait, multi lactation, single breed model. Test day records since 1988)
- Protein yield for Holstein is investigated
  - P-index, relative index
    - Base = 100, cows born in 1990 and 1991
    - STD = 10, genetic std. app. 10.5



## Material

- Data is from February 2012
- Results has been presented at Interbull meeting in Cork 2012



## Senarios

- Rout: Routine evaluation with all animals included
- Y 1990-2003: Pre-selection of young bulls born 1990-2003:
- Al bulls with progeny test above average selected Selection intensity 0.8, selection differential 0.75 Records from daughters of discarded bulls were set to missing (no progeny test)
- Y 2000-: Same as in previous scenario but for young bulls born 2000 and onwards





## Senarios (continued)

- Sec\_1995-: No pre-selection for young bulls, but second crop daughters born in 1995 and onwards have their records set to missing
- Y\_Sec\_2000-: Daughters born year 2000 and onwards :
  - No second crop daughters with records
  - Pre-selection of young bulls above average
- Daughter records were set to missing but pedigree information kept

All senarious: No changes for private and foreign bulls



## No of progeny tested bulls

Birth year	Rout	Y_1990-03	Y_2000-	Sec_1995-	Y_Sec_2000-
1990	412	<u>201</u>	412	412	412
1992	537	<u>269</u>	537	537	537
1994	469	<u>238</u>	469	469	469
1996	475	<u>236</u>	475	475	475
1998	450	<u>225</u>	450	450	<u>225</u>
2000	392	<u>182</u>	<u>182</u>	392	<u>182</u>
2002	398	<u>200</u>	<u>200</u>	398	<u>200</u>
2004	357	357	<u>181</u>	357	<u>181</u>
2006	393	393	<u>186</u>	393	<u>186</u>





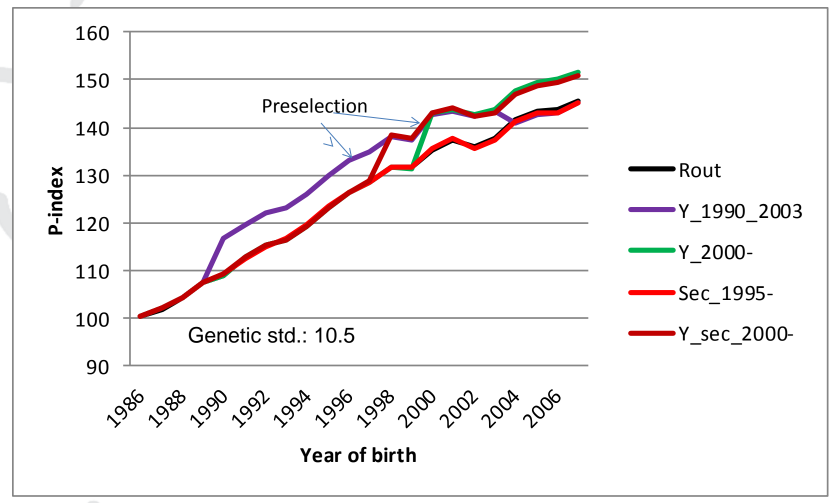
### Largest progeny group size

Birth year	Rout	Y_1900-03	Y_2000-	Sec_1995-	Y_Sec_2000-
1990	23654	23654	23654	<u>312</u>	20477
1992	54491	54491	54491	<u>350</u>	<u>25842</u>
1994	22013	22013	22013	<u>297</u>	<u>1644</u>
1996	53705	53705	53705	<u>244</u>	<u>244</u>
1998	37655	37655	37655	<u>396</u>	<u>396</u>
2000	21794	21794	21794	<u>216</u>	<u>214</u>
2002	20251	20251	20251	<u>227</u>	<u>212</u>
2004	552	552	552	<u>275</u>	<u>275</u>
2006	405	405	316	<u>405</u>	<u>316</u>





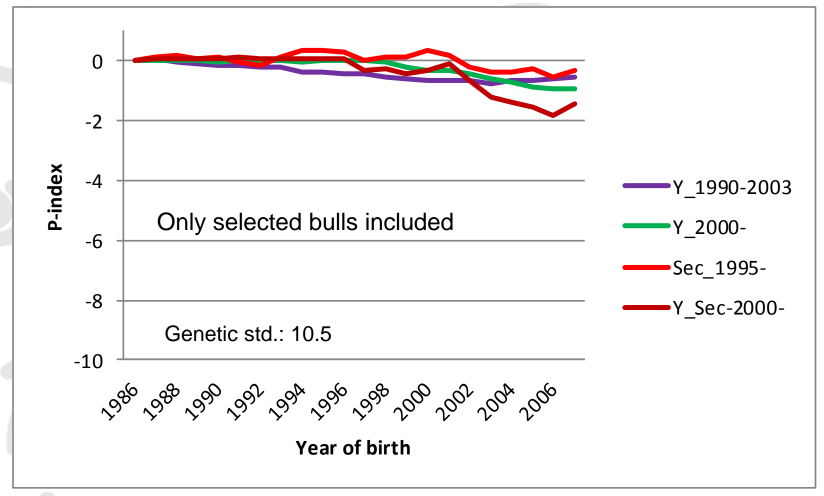
#### Genetic trend protein Nordic A.I. bulls





Different no of bulls in trend estimation

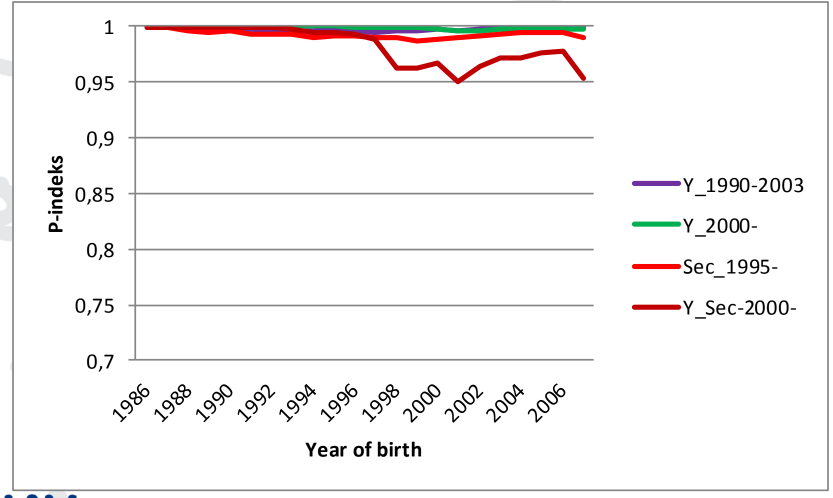
### Difference to routine run, protein







#### Correlation with routine run







#### Difference in P-index in scenario Y\_Sec\_2000- compared to routine run for bulls born after 2000. No correction for differences in trend

Difference	Frequency	Percent
-6	5	0.4
-5	5	0.4
-4	21	1.8
-3	155	12.9
-2	298	24.9
-1	348	29.1
0	223	18.6
1	104	8.7
2	33	2.8
3	6	0.5





#### Conclusion

- Pre-selection of young bulls with a reliability of 0.90 and a selection intensity of 0.8 has negative effect on the genetic trend
  - Bias in expected direction, i.e. trend is underestimated
  - Effect less than was expected
- Omitting second crop daughters has less effect on trend.
- Pre-selection of young bulls in combination with omitting second crop daughters has bigger but still minor effect
- Likely: Genomic selected bulls will be used over a longer time span than young bulls in the old young bull system, and some genomic tested bulls are used for nearly a year
- Selection intensity for a single trait is lower than for the Total Merit Index
- Genomic selection will not ruin traditional estimation of breeding values in the first years



