





Forensic science technique applied for calculation of kinship index

E. Bömcke 1,2, and N. Gengler 1,3

- ¹ Animal Science Unit, Gembloux Agro-Bio Tech, University of Liege (GxABT, ULg) – Gembloux, Belgium ² FRIA - Brussels, Belgium
- ³ National Fund for Scientific Research (FRS-FNRS) Brussels, Belgium

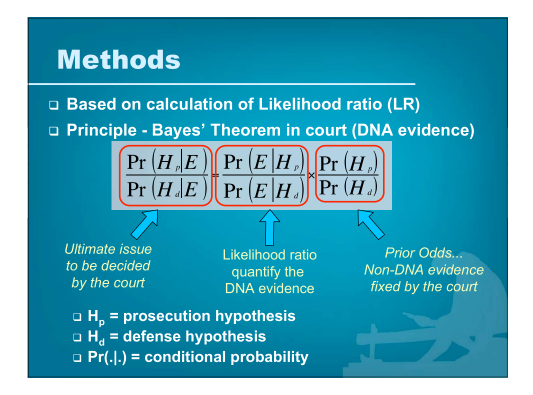
EAAP 2010 August 23 - August 27, Heraklion, Greece

Context

- □ Relationship coefficients = useful tool to improve genetic management of small and/or endangered populations
- □ Relationship coefficients are traditionally calculated on pedigree data
- □ But in pedigrees
 - □ Often presence of errors
 - □ Some parents are missing

Objective

- □ Find tools to help the breeders for the management of endangered population or population with incomplete pedigrees
- Increase the knowledge of kinship trough detection in the pedigrees of
 - □ False parents
 - □ Non-recorded parents
 - → Calculation of kinship index





□ Bayes' Theorem in court - Prosecution hypothesis H_p

$$\frac{\Pr(H_{p}|E)}{\Pr(H_{d}|E)} = \frac{\Pr(E|H_{p})}{\Pr(E|H_{d})} \times \frac{\Pr(H_{p})}{\Pr(H_{d})}$$

 $Pr(E|H_p)$ = probability that Mr X's DNA profile matches the crime profile given Mr X is the source of the crime profile

Mr X. is the killer



Principle

□ Bayes' Theorem in court - Defense hypothesis H_p

$$\frac{\Pr(H_{p}|E)}{\Pr(H_{d}|E)} = \frac{\Pr(E|H_{p})}{\Pr(E|H_{d})} \times \frac{\Pr(H_{p})}{\Pr(H_{d})}$$

Pr(E|H_d) = probability that Mr X's DNA profile matches the crime profile given Mr X is <u>not</u> the source of the crime profile <u>and that it has</u> <u>originated from an unrelated individual</u>

Mr X. is the killer



Another individual is the killer

Principle

□ Reporting the DNA evidence

LR	Support for H _p	LR	Support for H _d
> 106	Extremely strong	< 0.1	Limited
> 10 ⁵	Very strong	< 0.01	Moderate
> 104	Strong	< 10 ⁻³	Moderately strong
> 10 ³	Moderately strong	< 10-4	Strong
> 100	Moderate	< 10 ⁻⁵	Very strong
> 10	Limited	< 10-6	Extremely strong

LR = 1 inconclusive

Methods

- □ In court, DNA evidence match directly to criminal profiles
- □ But, LR can also be used to do Familial searching

Methods

- □ In court, DNA evidence match directly to criminal profiles
- □ But, LR can also be used to do Familial searching
- □ Familial searching
 - □ Method used in forensic science
 - □ Search for people in a database who are related to DNA evidence
 - □ New hypothesis for parentage calculations
 - □ H_p → H1: the alleged father (or mother) is the true parent
 - \Box H_d \rightarrow H2: the alleged father (or mother) is not the parent

Methods - parent/child LR

□ 2 individuals (x and y), 1 locus, 4 allele positions

$$LR = \frac{Pr(ab,cd|H_1)}{Pr(ab,cd|H_2)} = U$$

□ U depends on the frequencies of shared alleles

$$U=(x_1+x_2+x_3+x_4)/4$$

- \Box Define x_1 , x_2 , x_3 , x_4 for the allele pairs ac, ad, bc, bd
- □ x_i=1/p_i if the two alleles are the same type
- □ x_i=0 if the 2 alleles are not the same type

Methods - parent/child LR

- \Box LR = $(x_1 + x_2 + x_3 + x_4)/4$
- □ 8 possible situations (for each locus)

Genotypes	LR
ii,ii	1/p _i
ii,ij and ij,ii	1/(2p _i)
ij,ij	$(p_i+p_j)/(4p_ip_j)$
ij,(il or ki)	1/(4p _i)

Genotypes	LR
ij,(jl or kj)	1/(4p _j)
ij,ij	1/(2p _j)
(ii or ij),kl	0
Missing value	1

Methods - parent/child LR

- \Box LR = $(x_1 + x_2 + x_3 + x_4)/4$
- □ 8 possible situations (for each locus)

Genotypes	LR
ii,ii	1/p _i
ii,ij and ij,ii	1/(2p _i)
ij,ij	$(p_i+p_j)/(4p_ip_j)$
ij,(il or ki)	1/(4p _i)

Genotypes	LR
ij,(jl or kj)	1/(4p _j)
ij,jj	1/(2p _j)
(ii or ij),kl	0
Missing value	1

- □ For a set of n loci: LR_{set} = LR_{L1} x LR_{L2} x LR_{L3} x ... x LR_{Ln}
 - □ If only one LR=0 → genotyping error → LR=1

Data simulation

- □ Pedigree:
 - □ 100 years of simulation (1907-2007)
 - □ 3 repetitions with 3 levels of inbreeding
 - □ Ped1: ~17%
 - □ Ped2: ~27%
 - □ Ped3: ~37%
- □ Genotypes:
 - □ 25 microsatellites with 3 to 15 alleles
 - □ Equal allele frequencies in founder population
 - □ 5 repetitions/complete pedigree (data1 to data5)

Results

□ Number of animal in the pedigree = database size

	Nb of animals	Nb of comparisons
Ped 1	1,134	1,285,956
Ped 2	761	579,121
Ped 3	771	594,441

□ Number of animal in the pedigree = database size

	Nb of animals	Nb of comparisons
Ped 1	1,134	1,285,956
Ped 2	761	579,121
Ped 3	771	594,441

- □ In order to decrease the number of comparisons, use of
 - □ 'Local' prior information = information about pairs of individuals (e.g. sex, birthyear)
 - □ 'Global' prior information = general knowledge about population structure (e.g. generation interval, sexual maturity)

Results

□ Number of comparisons with prior information (PI)

	Nb of animals Nb of comparisons		Nb with PI
Ped 1	1,134	1,285,956	22,558
Ped 2	761	579,121	11,330
Ped 3	771	594,441	39,543

→ Reduction > to 90%, dependent of the inbreeding level of the pedigree

□ Number of comparisons with LR > 0 = possible parents

	Nb of animals	Nb with PI	Nb with LR > 0
Ped 1	1,134	22,558	307
Ped 2	761	11,330	334
Ped 3	771	39,543	2,815

→ Up to 98 % of the calculated parent-child combination are rejected (LR = 0), dependent from inbreeding

Results

□ Ranking of true parents (%)

	Ped1 (F	≈ 17%)	Ped2 (F	F≈ 27%) Ped3 (F≈ 37%)		
	Case A ¹	Case B ²	Case A ¹	Case B ²	Case A ¹	Case B ²
In first position	49.26	49.68	48.44	49.53	42.75	44.94
In 1 st and 2 nd position	97.16	96.95	90.94	92.03	73.71	73.31
In first 5 positions	100.0	100.0	99.69	99.84	93.93	93.76
In first 10 positions			100.0	100.0	98.26	98.88
In first 40 positions	7				100.0	100.0

¹ Case A: allele frequencies = frequencies from base population ² Case B: allele frequencies = frequencies from genotyped population

□ Ranking of true parents (%)

	Ped1 (F ≈ 17%)		Ped2 (F	≈ 27%)	Ped3 (F	Ped3 (F ≈ 37%)	
	Case A ¹	Case B ²	Case A ¹	Case B ²	Case A ¹	Case B ²	
In first position	49.26	49.68	48.44	49.53	42.75	44.94	
In 1 st and 2 nd position	97.16	96.95	90.94	92.03	73.71	73.31	
In first 5 positions	100.0	100.0	99.69	99.84	93.93	93.76	
In first 10 positions			100.0	100.0	98.26	98.88	
positions	→ High impact of inbreeding Highly inbred parents are more often					100.0	
	miss-ranked: full- (half-) sibs of the (grand-)parents are in higher position					1	

Results

□ Mean values of LR

	Ped1 (F ≈ 17%)		Ped2 (F ≈ 27%)		Ped3 (F ≈ 37%)	
	Case A	Case B	Case A	Case B	Case A	Case B
In first position	4.68E15	2.43E8	1.08E17	7.19E6	4.53E18	3.07E6
In 1 st and 2 nd position	2.67E15	1.24E8	6.50E16	3.66E6	2.68E18	1.54E6
In first 5 positions	1.41E15	6.23E7	2.84E16	1.47E6	1.24E18	6.18E5
In first 10 positions			2.04E16	1.22E6	7.58E17	3.09E5
In first 40 positions					3.38E17	1.54E5
Max and	1.33E17	5.27E10	2.13E18	5.97E8	5.46E20	2.10E9
min values	2.54E11	72.22	3.25E13	17.92	1.04E15	0.272

□ Mean values of LR

	Ped1 (F ≈ 17%)		Ped2 (F ≈ 27%)		Ped3 (F ≈ 37%)	
	Case A	Case B	Case A	Case B	Case A	Case B
In first position	4.68E15	2.43E8	1.08E17	7.19E6	4.53E18	3.07E6
In 1 st and 2 nd position	2.67E15	1.24E8	6.50E16	3.66E6	2.68E18	1.54E6
In first 5 positions	→ High impact of allele frequencies Disequilibrium in allele frequencies (rare vs very common alleles) decreases the mean value of LR and increases the range of values Inbred parents, carrying common alleles have very low LR values = risk to reject true parentage					
In first 10 positions						
In first 40 positions						
Max and min values	1.33E17	5.27E10	2.13E18	5.97E8	5.46E20	2.10E9
	2.54E11	72.22	3.25E13	17.92	1.04E15	0.272

Application to real data

- □ Skyros pony: an endangered Greek horse breed
 - □ Population size: about 200 individuals
- □ Available data for the breed
 - □ Partial pedigree
 - □ Total of 395 individuals
 - □ Pedigree deepness: 1.5 generation-equivalents
 - □ Genotypes of half of the living population (99 ind.)









- □ Skyros pony 99 individuals genotyped with
 - □ 2 parents genotyped (14 ind.)
 - □ Dam genotyped (28 ind.)
- → 62 parents
- □ Sire genotyped (6 ind.)
- □ Application of Familial Searching
 - □ 1 genotyping error max

- □ Ranking if 2 parents genotyped (14 cases)
 - □ 4 cases: sire and dam in positions 1 and 2
 - □ 4 cases: sire and dam in positions 1 and 3
 - □ 3 cases: sire and dam in positions 2 and 3
 - □ 3 cases: sire and dam in positions 1 and 5, 6 or 8

With for 5 parents 1 genotyping error

- □ Ranking if 2 parents genotyped (14 cases)
 - □ 4 cases: sire and dam in positions 1 and 2
 - □ 4 cases: sire and dam in positions 1 and 3
 - □ 3 cases: sire and dam in positions 2 and 3
 - □ 3 cases: sire and dam in positions 1 and 5, 6 or 8

With for 5 parents 1 genotyping error

→ High impact of inbreeding:

For most inbred individuals, possibility to have in first positions half-/full-sibs of the parents or grand-parents

Results

- □ Ranking if 1 parent genotyped (34 cases)
 - □ 22 cases: parent in position 1
 - □ 6 cases: parent in position 2
 - □ 2 cases: parent in position 3

With for 10 parents 1 genotyping error

- □ 4 cases: parentage rejected (LR=0)
 - □ In 3 cases, the true parent was detected in the genotyped set

→ High impact of

inbreeding

One supplementary parentage was detected in the set

Conclusions

- □ Familial searching offers promising results for improvement of pedigrees
 - → Detection of non-recorded or false parentages
- □ Efficiency of programs can be increased using prior information
- □ Ranking / LR values of true parents influenced by
 - □ Inbreeding
 - □ Allele frequencies used to calculate LR
 - □ Presence of genotyping errors / missing values

Corresponding author's email: elisabeth.bomcke@ulg.ac.be

Thank you for your attention!

Study supported by:

National Fund for Scientific Research (FRS – FNRS)

