

ISFG 2024 Workshop



Lecture 4. Disaster victim identification



Thore Egeland Norwegian University of Life Sciences (NMBU) & Department of Forensic Medicine, Norway

Disaster Victim Identification (DVI)

• DVI

- Match list of unidentified persons against a list of missing persons
- Special case:
 - Missing Person Identification



Cases

- Missing grandchildren of Argentina
- World Trade Center attack
- Spitsbergen civil aircraft disaster
- Balkan conflicts
- Drowned immigrants
- Thailand tsunami
- Methods and software
 - Exclusion and inclusion power
 - Treat *each* victim (PM approach) or *each* family (AM approach) at a time
 - Joint approach. GLR
 - library(dvir)



First: Missing Person Identification



- Special case of DVI:
 - ✓ Only one *missing person* (MP) in each reference family
 - ✓ Possibly many reference families and many *persons of interest* (POIs)
 - ✓ Each case treated independently

Exclusion power



- Sufficient data to *exclude* an unrelated POI?
- Exclusion power

EP = P(data incompat. with ped | POI unrelated)

 Can be computed exactly (Egeland, Pinto, Vigeland, 2014). In forrel: > exclusionPower() > missingPersonEP()

Inclusion power



- Sufficient data to give a positive match if POI = MP?
- **Inclusion power** (or *exceedance probability*):

 $IP_{10000} = P(LR > 10000 | POI = MP)$

- Can be computed by simulation!
 - Unconditional \rightarrow <u>average</u> for all pedigrees of this type
 - Conditional \rightarrow probability for <u>this particular case</u>

In forrel: > missingPersonIP()



Argentina 1976 – 1983: Dirty War

- Military dictatorship
- War against guerrillas

• Opponents killed or disappeared

- About 500 children abducted
 - kidnapped with their parents or born in captivity
 - parents killed
 - raised by police or military families







History

- 1977: Grandmothers of Plaza de Mayo
 - weekly marches ever since
 - <u>https://www.abuelas.org.ar</u>
- 1983: Mary-Claire King, US geneticist
 - Collaboration with the Grandmothers
 - HLA typing + blood groups
 - First grandchild recovered using genetics
- 1987: National Genetic Data Bank (BNDG)
- 2007: Argentine Forensic Anthropology Team
- 2021: In total 130 reunifications by BNDG





BANCO NACIONAL DE DATOS GENÉTICOS

Exclusion and inclusion



- POI1 = 12/12 is excluded if mutations are ignored since the alleles of his father are among 9/10/13
- POI2 is
 - $\checkmark \quad \text{not excluded}$
 - ✓ included if LR > 10000

Power analysis

Power evaluation of ~200 families in the BNDG database
most of them unsolved

Typed	Families	Parent(s) typed	2 nd degree only		
1	11	5	5		
2	20	5	15		
3	29	7	20		
4	41	6	33 14 14		
5	29	7			
6	26	5			
7	14	0	8		
8	14	0	9		
9	5	0	3		
10+	7	0	5		
Total	196	35	126		

Missing	Total	mtDNA	Y
Male	27	27	25
Female	17	17	-
Unknown	152	142	108
Total	196	186	133

• For each family: Compute IP₁₀₀₀₀ and EP



Disaster victim identification (DVI)



Initial steps

- Merge identical samples
- Direct matching
- Remove non-identifiable
- ➤ Exclude

Pairwise	ІВ Н.	$V_i = M$	ivs H	·unrelated
rall wise	LЛ. П ₁ .	$v_1 - v_1$	ј vs. п ₂	. umerateu

	M1	M2	M3
V1	1	0	0.39
V2	1	16142	0
V3	1	0	6266

Challenge: Sometimes pairwise fails



	М1	M2	МЗ
V1	1	20	10.5
V2	1	0	0.5
V3	1	10	5.5

Pairwise LRs indicate V1 = M2, but

• V1 = M2 is impossible if V2 or V3 are among the missing

Joint solution



Joint likelihoods, sorted

	V1	V2	V3	loglik	LR
1	M3	M1	M2	-15.67181	2000.0
2	M2	*	M1	-17.97439	200.0
3	M2	*	M3	-17.97439	200.0
4	*	M1	M2	-17.97439	200.0
5	M3	*	M2	-18.66754	100.0
6	*	M3	M2	-18.66754	100.0
19	*	*	*	-23.27271	1.0
34	*	M3	*	-Inf	0.0

- Two problems:
 - ✓ LR uses V1 = '*', V2 = '*', V3 = '*' as comparison
 - ✓ How do we test e.g. H_1 : V1 = M3 vs. H_2 : V1 = '*'?

A tool for interpreting joint results: Generalised likelihood ratio (GLR)

 $GLR = \frac{\max_{i \in H_1} L_i}{\max_{j \in H_2} L_j}$



	V1	V2	V3	loglik	LR
1	M3	M1	M2	-15.67181	2000.0
2	M2	*	M1	-17.97439	200.0
3	M2	*	M3	-17.97439	200.0
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5	М3	*	M2	-18.66754	100.0
6	*	М3	M2	-18.66754	100.0
19	*	*	*	-23.27271	1.0
					•••
34	*	M3	*	-Inf	0.0

$$\mathrm{GLR} = \frac{LR_1}{LR_4} = \frac{2000}{200} = 10$$

Interpretation of GLR = x

 $\text{GLR} = \frac{\max_{i \in H_1} L_i}{\max_{j \in H_2} L_j}$

The best explanation of the data given H1 is x times more likely than the best explanation given H2

Symmetric matches

				Joint likelihoods				
PM	AM	Question 2:		V1	V2	V3	loglik	_
			1	M1	M2	M3	-257.7	v
			2	M1	M3	M2	-257.7	v
V1		$\{1/1, 1/2\} = \{1/1, 1/2\}$	3	*	M2	M3	-268.9	v
		[VI, VZ] = [IVIZ, IVIS]:	4	*	M3	M2	-268.9	v
			5	M2	M1	M3	-272.6	د
V2	R1 M1		6	M3	M1	M2	-272.6	۲
			7	M1	*	M2	-276.5	X
	? ~ ())		8	M1	*	M3	-276.5	۲
V3	M2 M3		9	M2	M3	M1	-276.6	د
L			:					
			32	*	*	*	-286.0	X
			33	*	M1	*	-286.5	5

M1 -292.2

x

*

34

*

$$GLR = 2.9 \times 10^{6}$$

strong evidence

Discussion

- What should be reported?
 - LR and, or posterior?
 - Thresholds?
 - Verbal scales?

Biased selection of references



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Joint DNA-based disaster victim identification

Magnus D. Vigeland 🗠 & Thore Egeland

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