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A comment on the Paper:

Extended Y chromosome haplotypes resolve multiple and unique lineages of the Jewish Priesthood

By M.F. Hammer, D.M. Behar, T.M. Karafet, F.L. Mendez, B. Hallmark, T. Erez, L.A. Zivotovsky, S. Rosset, K. Skorecki, Hum. Genet., Published online 08 August 2009

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Letters to the Editors

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The cited paper is remarkable with respect to haplogroups and haplotypes typing and the large number of individuals tested, including Cohanim, which is the prime target of the study. However, it is open to critique regarding calculations of time spans to common ancestors of presented series of haplotypes.

First, the authors have employed an inadequate methodology for the calculations, namely the infamous "population mutation rates" of 0.00069 mutation/marker/generation. It should not have been used in the first place. Much more adequate procedures have been developed lately. Second, the authors have not defined criteria when the "population" and when the "pedigree", or the "genealogical" mutation rates, should be used. As a result, they have applied the "population" mutation rate to "genealogical" haplotype series, which increased their "TMRCA", that is a time span to common ancestors, by about 300%. Third, they have not analyzed genealogical lineages with recent common ancestors (such as 500-800 years bp), therefore, they missed valuable information regarding history of Cohanim in this millennium.

Allow me to explain.

Cohanim J1e*-P58*

Here is the haplotypes tree (it is explained in the last section of this Comment) for Cohanim, haplogroup J1e*, composed from the Hammer et al (2009) data, 22-marker haplotypes, Table S3.

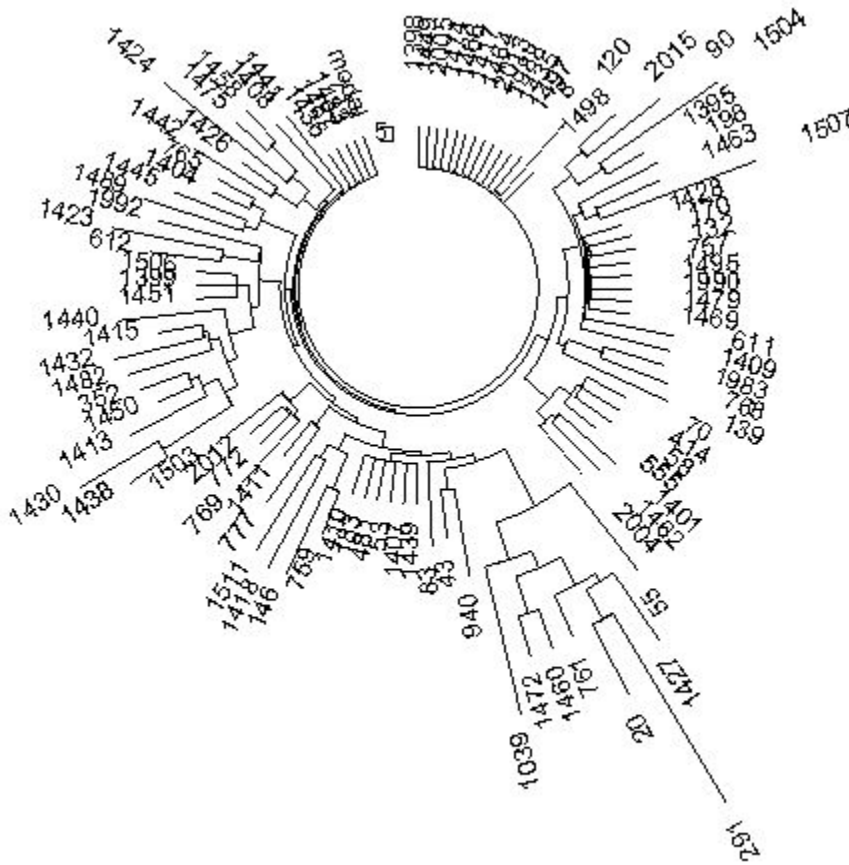


Figure 1. 22-marker haplotype tree for Cohanim, haplogroup J1e*-P58*, for 98 haplotypes (Hammer et al, 2009). Haplotypes numbering corresponds to the numbering in the paper, Table S3

The tree contains 16 base (ancestral) haplotypes (see the last section for definitions) on the top (in the entire order DYS 393, 390, 19, 391, 385a, 385b, 426, 388, 439, 389-1, 393, 389-2, 458, 459a, 459b, 455, 454, 447, 437, 448, 449 and 438):

12 23 14 10 13 15 11 16 12 13 11 30 – 17 8 9 11 11 26 14 21 26 10

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3 All 98 haplotypes contain 191 mutations from the above base haplotype, which
4 gives – employing the linear method (see the last section of this paper) –
5 $191/98/0.047 = 41$ generations (without a corrections for back mutations
6 [Klyosov, 2009]), or 43 generations with the correction, to a common ancestor,
7 that is 1075 ± 130 years bp, the 10-th century AD plus-minus a century.
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11 The authors of the cited paper have obtained 3200 ± 1100 years to a common
12 ancestor for the Cohanim, using the same haplotypes, albeit with a set of 17-
13 marker haplotypes, and 3000 ± 1500 years to a common ancestor using a set of 9-
14 marker haplotypes, and claimed that is it nicely fit to the Biblical description for
15 the origin of the Jewish higher priest. It is not. See below for more details.
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19 Since all 98 haplotypes contain 16 base (identical to each other) 22-marker
20 haplotypes (see above), it gives – employing the logarithmic method (see the last
21 section of this paper) – $\ln(98/16)/0.047 = 39$ generations (without a correction),
22 or 41 generation (with the correction), that is 1025 years bp to a common ancestor
23 for Cohanim J1e* haplotypes.
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27 Since the logarithmic and the “linear” methods give the same dating to the
28 common ancestor, it means that there was indeed just one common ancestor for
29 the whole series of 98 of 22-marker haplotypes (see the last section), who lived
30 about 1075 ± 130 years bp.
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33 The authors of the cited paper have removed some markers for their analysis,
34 namely five markers, DYS 385a, 385b, 459a, 459b and 458, reducing the series
35 from 22 to 17-marker haplotypes. Let us consider if there would be any
36 noticeable change. 98 of 17-marker haplotypes contain 140 mutations from the
37 above base haplotype (less of the five markers), which gives 1175 ± 160 years (bp)
38 for the common ancestor. It is practically the same timespan within the margin of
39 error.
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43 As it was noticed above, the authors of the cited paper have obtained 3200 ± 1100
44 years to a common ancestor for the Cohanim, using the same set of 17-marker
45 haplotypes, and claimed that is nicely fits to the Biblical description for the origin
46 of the Jewish higher priest. It is not, with 1075 ± 130 and 1175 ± 160 years (bp).
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49 Why such a large difference? The answer is rather obvious. The ratio of 3200 and
50 1075 years is 3.0. The ratio of the “population” mutation rate (0.00069) to the
51 actual, “genealogical” one (0.00214) is 3.1. The authors of the cited paper have
52 applied the “population” mutation rate to the “genealogical” series of
53 haplotypes. Actually, that “population” mutation rate should have never been
54 applied in the first place. It is ill defined, it is not governed by any criteria when
55 it should be applied and when it should not.
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In Table S4 of the cited paper the authors listed 99 of 12-marker Cohanim J1e* (J-P58*) haplotypes. All of them contain 98 mutations from the above base haplotype (in the first 12 markers), which gives 1175 ± 170 years to a common ancestor. It is the same figure obtained above, within the margin of error.

Finally, the authors reduced the 22-marker haplotypes to 9-marker one (the 12-marker haplotype in the FTDNA format less of DYS 385a, 385b and 426), and, using the same "population" mutation rate, have obtained 3000 ± 1500 years to a common ancestor of Cohanim J1e*. In fact, this whole series contain 67 mutations, which gives $67/98/0.018 = 38$ generations (40 generations with a correction for back mutations). That is 1000 ± 160 years to a common ancestor. This is again practically the same figure as given above for 22-marker (1075 ± 130 ybp), 17-marker (1175 ± 160 лет), and 12-marker (1175 ± 170 ybp) haplotypes, and is far away from the authors' 3000 ± 1500 ybp for Cohanim J1e*.

In fact, the ancestral haplotype of Cohanim J1e* (in the format previously described)

12 23 14 10 13 15 11 16 12 13 11 30 – 17 8 9 11 11 26 14 21 26 10

was identified in (Klyosov, 2008a; 2009b) in the form of a 67-marker haplotype (the respective 22 markers are shown below for a comparison)

12 23 14 10 13 15 11 16 12 13 11 30 -- 17 8 9 11 11 26 14 21 27 10

(so-called "the recent extended CMH") of a common ancestor who lived $1,050 \pm 190$ years bp, around the 10th century. One can see, it is practically the same ancestral haplotype and the same time span to a common ancestor.

Recent DNA-genealogical branches of Cohanim J1e*

The tree (Fig. 1) reveals three principal branches (on top and the left-hand side; on the right-hand side; and at the bottom), with their ancestral haplotypes, respectively :

12 23 14 10 13 15 11 16 12 13 11 30 – 17 8 9 11 11 26 14 21 26 10

12 23 14 10 13 15 11 16 12 13 11 30 – 17 8 9 11 11 26 14 21 27 10

12 23 14 10 13 15 11 16 **11** 13 11 30 – 17 8 9 11 11 26 14 21 26 10

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3 That the “young” branches are really young ones can be seen from Fig. 2, which
4 shows a haplotype tree for 12-marker haplotypes of the Cohanim of haplogroup
5 J1e*, listed in the cited paper.
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9 Since these haplotypes are lacking ten markers after their first twelve ones,
10 compared to those in Fig. 1, the tree is slightly rearranged in Fig. 2, following the
11 computer program. However, it shows the main tree branches in the haplotype
12 series. Since the branches lost many mutations (in markers 13 through 22), “ages”
13 of their common ancestors are reduced for some branches. For example, the
14 right-hand side branch now contains 48 haplotypes, 43 of which are identical to
15 each other (base haplotypes), and the other contains only seven mutations. This
16 gives $\ln(48/43)/0.022 = 5.0$ generations, that is 125 ± 50 years to a common
17 ancestor, and $7/48/0.022 = 6.6$ generations, that is 165 ± 65 years to a common
18 ancestor. Since the logarithmic and linear methods again give practically
19 identical results, it points to a single (in terms of DNA genealogy) common
20 ancestor for the whole branch. It also shows that 12-marker haplotypes contain
21 much poorer information regarding a time span to a common ancestor compared
22 to that of 22-marker haplotypes. Hence, it is surprising why the authors were
23 reluctant to consider the 22-marker haplotypes of Cohanim, persistently
24 reducing a number of markers in the haplotypes to 17, 12, and even 9.
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30 Cohanim J2a-M410*. 31

32 The 22-marker haplotype tree is shown in Fig. 3
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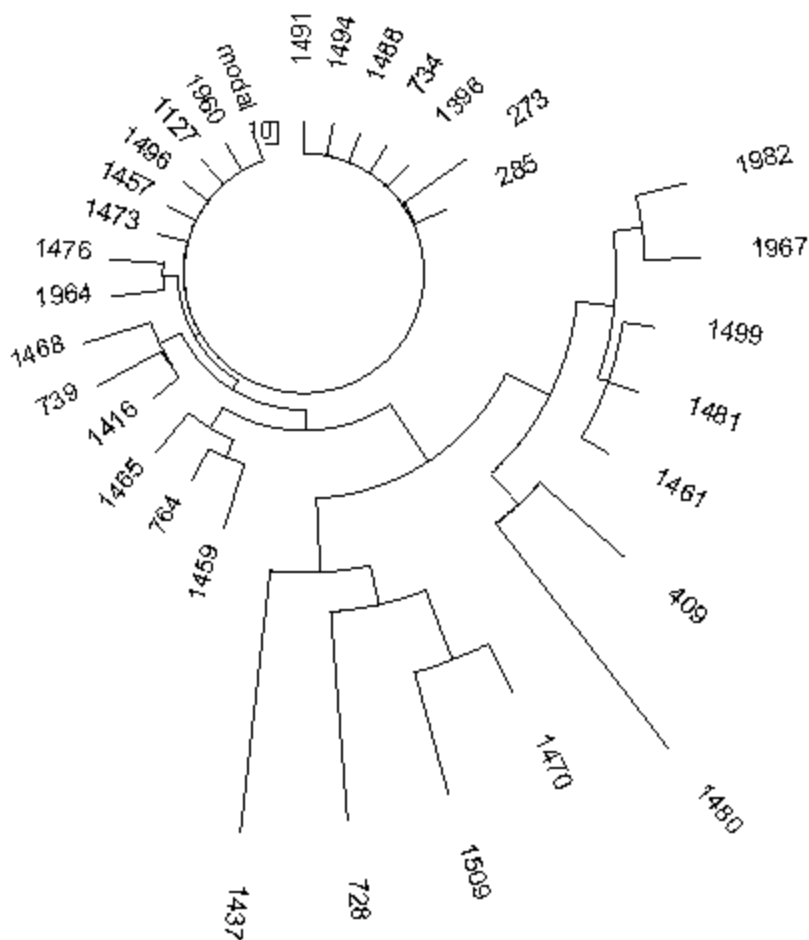


Figure 3. 22-marker haplotype tree for Cohanim, haplogroup J2a-M410, for 31 haplotypes (Hammer et al, 2009). Haplotypes numbering corresponds to the numbering in the paper, Table S3

One can see that the tree consists of two quite different branches, the older one in turn contains two sub-branches. It is clear that to treat the tree as a whole, without subdivision into branches, as it is done in the cited paper, would be incorrect and lead to a phantom common ancestor (though the authors in the paper actually indicated this may be a problem as not all SNPs down stream from M410 were considered). However, the authors did exactly that, they took the whole series of 31 haplotypes, applied the “population” mutation rate and obtained 5900 ± 2000 years to a common ancestor using 17-marker haplotypes, and 4900 ± 1900 years to a common ancestor using 9-marker ones. Again, they have eliminated markers DYS385a,b and DYS459a,b, as well as DYS449, and reduced the 22-marker haplotypes to 17-marker ones. It is obvious that the younger series of haplotypes and the respective recent common ancestor were lost from considerations in the cited paper.

Let us consider the younger branch, containing 20 of 22-marker haplotypes, having 14 mutations from the base haplotype (in the format described above):

12 23 15 10 14 17 11 16 12 14 11 30 – 15 8 9 11 11 25 15 21 31 9

It contains 11 base haplotypes, hence, $\ln(20/11)/0.047 = 13$ generations to a common ancestor. The linear method gives $14/20/0.047 = 15 \pm 4$ generations to a common ancestor, which is practically the same value obtained by the logarithmic method, within margin of error. Therefore, a common ancestor for the “recent” Cohanim branch lived 375 ± 110 years bp, around the 17th century AD.

The older branch split to two sub-branches. The first one contains five haplotypes of descendants of Cohanim of the former Russian Empire, namely Russia, Ukraine, Belarus, Lithuania and Poland, with only three mutations per 110 markers from the base haplotype:

12 23 15 10 14 17 11 **15** 12 **13** 11 29 – 15 8 9 11 11 **24** 15 21 **32** 9

which gives 325 ± 190 years to their common ancestor, who lived around the 17th century. The alleles in bold are different from ones of the above base haplotype with a common ancestor of 375 ± 110 ybp. This places their common ancestor to about 1500 ybp, around the 6th century AD.

It is exactly the ancestral haplotype of Jewish J2 population identified in a 37-marker format (Klyosov, 2008a) (the respected alleles are shown here for a comparison):

12 23 15 10 14 17 11 15 12 13 11 29 – 15 8 9 11 11 24 15 21 32 9

with a common ancestor who lived 850 ± 260 years bp (12 mutations in 148 markers).

The older sub-branch also splits into two sub-branches, one includes descendants from Iran, Iraq, Argentina and North Africa, with a common ancestor of 3650 ± 830 years bp (24 mutations in four 22-marker haplotypes). The last two branches (Cohanim from the Russian Empire, on the one hand, and Iraq, Iran, Argentina and Africa, on the other) differ by 12 mutations per 22 markers, that places their common ancestor to about 6200 ybp.

If, following the authors of the cited paper, to reduce the amount of markers from 22 to 17, then the “young” branch contains 9 mutations, and a common

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3 ancestor lived $9/20/17/0.00184 = 14$ generations ago, that is 350 ± 120 ybp. This is
4 practically the same as 375 ± 110 ybp, determined using 22-marker haplotypes.
5 The Russian Empire Cohanim (see above), whose common ancestor lived
6 325 ± 190 ybp (22-marker haplotypes), now fits the value of 325 ± 230 ybp (17-
7 marker haplotypes, two mutations in 110 markers), that is just the same value. It
8 is not clear why the authors of the cited paper reduced the haplotypes from 22-
9 marker to 17-marker, 12-marker and 9-marker haplotypes. 22-marker ones
10 worked perfectly.
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15 The same was for the ancient branch. 24 mutations in four 22-marker haplotypes
16 (3650 ± 830 ybp) reduced to 16 mutations in four 17-marker haplotypes (3575 ± 960
17 ybp), which is just the same, albeit with a larger margin of error.
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20 For 9-marker haplotypes the “younger” branch of 20 haplotypes contains 9
21 mutations (625 ± 220 ybp) which for the 22-marker haplotypes it contains 14
22 mutations (375 ± 110 ybp). These values are similar and fit within margin of error,
23 and at any rate point at a recent common ancestor of Cohanim J2a.
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26 Finally, for ancient Cohanim J2a branches, 22- and 9-marker haplotypes give
27 similar time spans to a common ancestor: in the first case 24 mutations in four
28 haplotypes results in 3650 ± 830 years to a common ancestor, in the second case
29 the number of mutations reduced to 7, and gave 2425 ± 1200 years to a common
30 ancestor. One can see that the two values are within the margin of error,
31 however, 9-marker haplotypes give much larger error margin.
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36 Cohanim J2e-M12

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38 Fifteen of 22-marker haplotypes of Cohanim J2e-M12 are listed in Table S3 in the
39 cited paper (the authors have reported in Table 1 that they have considered 16
40 haplotypes). According to the cited paper, a common ancestor of the haplotype
41 lived 12100 ± 4400 years bp as determined with 22-marker haplotypes. The
42 haplotype tree is shown in Fig. 4.
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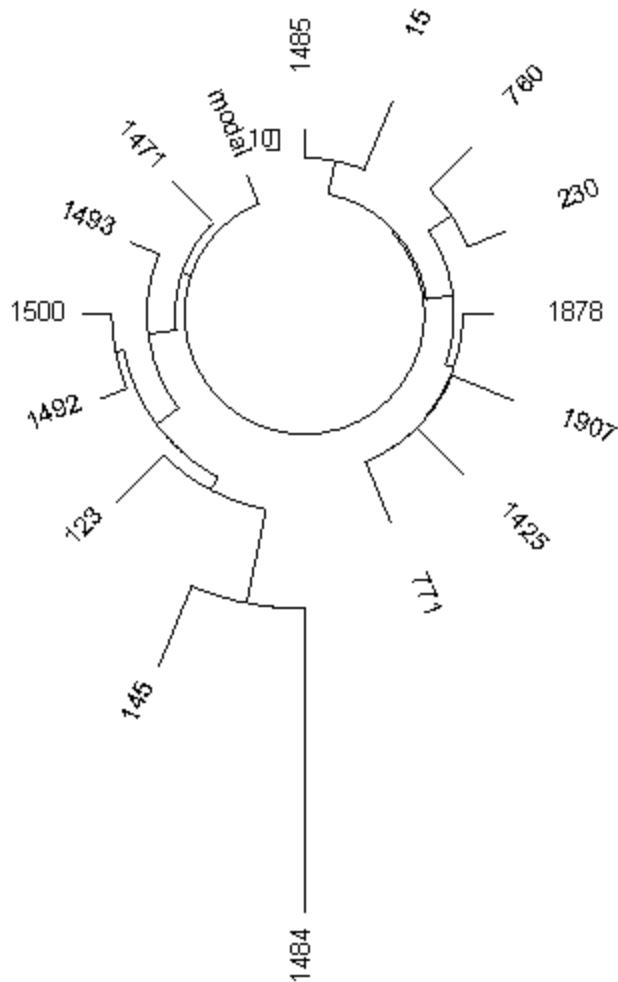


Figure 4. 22-marker haplotype tree for Cohanim, haplogroup J2b-M12, for 15 haplotypes (Hammer et al, 2009). Haplotypes numbering corresponds to the numbering in the paper, Table S3

The tree reveals two branches. The 8-haplotype branch on the right-hand side contains 15 mutations from the following base haplotype (in the format, explained above)

13 24 15 10 15 17 11 15 12 12 11 29 – 19 8 9 8 11 27 16 19 29 9

It gives $15/8/0.047 = 40$ generations (not corrected for back mutations), or 42 generations (corrected), that is 1050 ± 290 years to a common ancestor.

The 7-haplotype branch on the left-hand side contains 37 mutations from the base haplotype

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12 24 15 10 15 17 11 15 12 12 11 28 – 18 8 9 8 11 27 16 19 29 9

It gives 3150 ± 600 years from a common ancestor of the branch. The two base haplotypes are separated by three mutations, which places THEIR common ancestor to about 2950 years bp, which is the common ancestor of the left-hand branch itself. In other words, the right-hand side branch is a daughter branch of the left-hand side one, and a common ancestor of the whole tree of haplotypes of Cohanim J2b-M12 lived 3150 ± 600 years bp.

The authors of the cited paper, in which the time to a common ancestor was determined as 12100 ± 4400 ybp, were in error. They have treated the “father” and the “daughter” branches of the haplotype series as a just a set of haplotypes with a common ancestor outside of the branches, and, besides of it, employed an erroneous “population” mutation rate, as described above. Then, they have considered 17-marker haplotypes, omitting DYS 385a,b, DYS459a,b, and DYS449. These 15 of 17-marker haplotypes in fact contained 9 and 29 mutations in the right-hand side and the left-hand side branch, respectively, which results in 900 ± 310 and 3725 ± 800 years to their common ancestors, respectively. These figures are essentially the same with those obtained with the 22-marker haplotypes, within the margins of error.

Finally, if to reduce haplotypes to 9-marker ones, which would greatly sacrifice accuracy of calculations, the respective time spans to their common ancestors would be 1100 ± 460 and 1500 ± 600 years to a common ancestor, respectively, for the right-hand and left-hand side branches. It just could not possibly be 5500 ± 1900 ybp, obtained in the cited paper with the 9-marker haplotypes. The main reason of the errors, again, is wrong methodology employed in the cited paper for calculations of time spans to common ancestors.

Conclusions

Cohanim J1e*-P58*

A common ancestor of all 99 Cohanim lived 1075 ± 130 years bp, and this timing is reproducible for 9-, 12-, 17-, 22- and 67-marker haplotypes. A much higher values of 3190 ± 1090 and 3000 ± 1500 years bp were obtained in the cited paper (Hammer et al, 2009) by using incorrect methods and incorrect mutation rates. A common ancestor of all the 99 J1e* Cohanim lived around the 10th century AD. There are three main lineages derived from the common ancestor, with their common ancestors who lived approximately between 625 and 875 years bp.

An emphasis of the cited paper on the conclusion that “an extended CMH on the J1e*-P58* background that ... is remarkably absent in non-Jews” and having “the

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3 estimated divergence time of this lineage... 3,190±1,090 years” is incorrect
4 regarding the divergence time. It is much more understandable why the lineage
5 originated only 1,075±130 years ago is “remarkably absent in non-Jews”.

8 **Cohanim J2a-M410***

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11 Common ancestors of 31 Cohanim lived 325±190 and 375±110 years bp (recent
12 different DNA-lineages), and their common ancestor lived around 1500 ybp. A
13 common ancestor of another branch of Cohanim of this haplogroup lived
14 3560±830 years bp, and yet another common ancestor lived about 6200 years bp.

17 **Cohanim J2b-M12**

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20 Common ancestors of 15 Cohanim lived 3150±600 and 1050±290 years bp, with
21 the second lineage being descendant from the first common ancestor.

25 **Analysis of mutations and their rates**

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28 Haplotype trees were constructed using PHYLIP, the Phylogeny Inference
29 Package program (Felsenstein, 2005), as it was explained in detail in (Klyosov,
30 2008b; 2009a). A “comb” around the wheel, a “trunk”, in haplotype trees
31 identifies “base” haplotypes, identical to each other and carrying no mutations
32 compared to their ancestral haplotypes. The farther the haplotypes lies from the
33 wheel, the more mutations they carry compared to the base haplotype and the
34 older the respective branch.

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38 The “base” haplotypes are the ancestral haplotypes in an ideal case. However,
39 since those haplotypes often are deduced ones, it would be inaccurate to call
40 them “ancestral”. Hence, “base” haplotypes.

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43 The timespan to a common ancestor were calculated using both “logarithmic”
44 and “linear” approaches. The logarithmic method is based on the assumption
45 (which largely, or practically always holds) that a transition of the base
46 haplotypes into mutated ones is described by the first-order kinetics:

$$49 \quad B = Ae^{kt},$$

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52 that is

$$54 \quad \ln(B/A) = kt$$

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57 where:

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3 B = a total number of haplotypes in a set,
4 A = a number of unchanged (identical, not mutated) base haplotypes in the set,
5 k = an average mutation rate,
6 t = a number of generations to a common ancestor.
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10 One can see that the logarithmic method does not consider mutations in
11 haplotypes; only mutated and non-mutated (base) haplotypes are considered.
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14 Mutation-counting methods are all based on accumulation of mutations in
15 haplotypes over time. They include the "linear", "quadratic" (ASD) and
16 "permutational" methods (Klyosov, 2009a). In this paper only "linear" method is
17 employed. In this method a total number of mutations in a set of haplotypes is
18 counted, an average number of mutations per marker is calculated, a correction
19 for back mutations is introduced, either numerically, or using a handy table
20 (Klyosov, 2008c; 2009a), and a time span to a common ancestor is calculated,
21 either using the Table (Klyosov, 2008c; 2009a), or applying the respective
22 mutation rates. In other words, it is described by the following equation
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$$n/N/\mu = t$$

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29 where n is a number of mutations in all N haplotypes in the given series of
30 haplotype, μ is an average mutation rate per haplotype per generation, and t is a
31 number of generations to a common ancestor, corrected for back mutations.
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34 Average mutation rates employed in this paper, are:
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36 For 22-marker haplotypes – 0.047 mutations per haplotype, 0.00214 mutations
37 per marker; this value was obtained by summing up all 22 mutations rates for
38 each marker, according to (Chandler, 2006).
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41 For 17-marker haplotypes – 0.032 mutations per haplotype, 0.00188 mutations
42 per marker; this value was obtained by summing up all 17 mutation rates for
43 each marker, according to (Chandler, 2006).
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46 For 12-marker haplotypes – 0.022 mutations per haplotype, 0.00183 mutations
47 per marker; this value was obtained by summing up all 12 mutation rates for
48 each marker, according to (Chandler, 2006).
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51 For 9-marker haplotypes – 0.018 mutations per haplotype, 0.00200 mutations per
52 marker; this value was obtained by summing up all 9 mutation rates for each
53 marker, according to (Chandler, 2006).
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3 Error margin for time spans to common ancestors are calculated as described in
4 (Klyosov, 2009a), and correspond to the 95% confidence interval.
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7 Corrections for back (reverse) mutations can be done either numerically
8 (Klyosov, 2009a), or using the conversion table (Klyosov, 2008c; 2009a)
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