ABO GENETIC SYSTEM, SEXUALLY TRANSMITTED INFECTIONS AND ANDROGEN-ASSOCIATED DERMATOSES

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In the middle of 20th century, it was proved that ABO genetic system was the result of play of selection, including the infectious mortality due to two deadly epidemic collisions in the antiquity and with the main scene being Asian societies. It was discovered that plague tended to kill blood group O while smallpox blood group A carriers. Onwards no link was sought between this evolutionary phenomenon and blood group-related sexually transmitted infections and recurrent androgen-associated dermatoses (such as pityriasis versicolor and acne vulgaris) as well as sexual and fertility activity. Here we Dance Round such possible links. We found that these are expressed more strongly by blood group B carriers, and an attempt at translation of some relationships into population (intercontinental) level. We emphasize the genesis of blood group-related population gene pool equilibrium level and its attributes such as complex defense responses and co-operated immune reactions. Biomed Rev 2011; 22: 77-80.

Key words: ABO genetic system, cell-mediated immunity, population gene pool, sexually transmitted infections, recurrent androgen-related dermatoses

INTRODUCTION

Apart from significant advances in the study of human sexual potency, many aspects remain to be elucidated. The difficulty is in the fact that such an essential biological function and sociocultural phenomena along with related sexual practices cannot be reduced to a common denominator. In the middle of 20th century, it has been established that blood group B (BG-B) is a factual marker of Eastern belonging. Blood group B distribution in India preserves approximately equal values: for 1942 – 34.8% (1), for 1966-1970 - 32-42% (2), and for 1997 – 37.4% (3). Values concerning Japan are even more stable: for 1933-1944 21.9-23.1% (1) and for 1966-1970 – 22.2% (2). Here the reasons consist in the emigration processes which do
not alter BO polymorphism. Conversely, for the British Isles (4) and Germany (2), BG-B incidence increases on the expense of BG-A because of the intensive post-war immigration.

**Blood groups and antibody responses towards some sexually transmitted infections and their tropism to some androgen-associated dermatoses**

Research on the relationship between blood groups and sexual potency continuously grow in depth and quantity (5). We were inspired to do this investigation when during the one-act mass screening seroepidemiologic testing in 2001 among a total of 536 young, 18-year-old Caucasian navy sailors. They were examined a year before joining the Navy and then every season, i.e. 12 times during their 3-year service as well as every two years after dismissal. Telephone interviews as well as air-mailed or e-mailed questionnaires were used.

The purpose of the study was to establish if some well-known recurrent androgen-associated dermatoses (RAAD) such as *pityriasis versicolor* (RPv) (when starting during navy or permanently, Pm) and *acne vulgaris* (RAv) were expressed more strongly by BG-B carriers. It was established that BG-B carriers were higher generators of elevated antibody titres (> 1/20) against sexually transmitted infections (STI) such as herpes simplex virus-2 (HSV-2) (p<0.05 - p<0.001) and cytomegalovirus (CMV) (p<0.001) (Table 1) than the other BG carriers. The diagnosis of CMV infection was made by a classic hemagglutinin test while that of HSV-2 by additional clinical monitoring concordant in more than 85% of the cases with antibody responses. The possible BG-B-related higher sexual potency was tested during a 10-year long (2000-2009) randomized longitudinal population follow-up study. In order to achieve a higher preciseness we examined under homogenous endogenous and exogenous conditions such as matched gender, age, secondary educational level, nutrition, military stress level, inhabitants in the coastal area as well as origin from classic population morphs (CPMs) i.e., hereditary villagers (HVs) and hereditary town dwellers (HTDs).

The cause for discriminating the town-village hybrids (TVHs) will be stated later on. The concept that ABO group polymorphism results from the play of the natural selection, i.e., the infectious mortality from plague and smallpox (2-4) goes with our data where BG-B dominates by 3.58 times among HTDs (n=107) (in 43%) than among VHs (n=429) (in 12%) (p<0.001) being a consequence of the naturally rare disposition of the latter to these fatal infections.

As shown on Table 1, these RAAD gravitated by 1.25-2.16 times and by 1.65-2.0 times (p<0.01 - p<0.001), respectively, more strongly towards BG-B carriers than towards the other BG ones. A fascinating example was the unique strong BG-B carriers’ tropism towards double (RPv and RAv) RAAD (p<0.05 - p<0.01). On the other hand, there was no similar correlation concerning the general recurrence rate of dandruff (Rdf). This is in accordance with the hypothesis that Df, an abridged version of seborrhoic dermatosis, is a particular kind of eczema aggravated by added commensal, *Mallassezia* yeast rather than an infection *sui generis* as considered by some authors (6,7).

Concerning the aggregations of RAAD among BG-B carriers such as RPv+RAv, RPv+Rdf, and RAv+Rdf it is evident that Df is closer to Pv than to Av, however, there is no correlation to the high antibody responses towards HSV2 and CMV. Apparently, epidemiodynamics of Pv directly correlates with BG-B population frequency rate. Our data from illustrate a statistically significant BG-B domination by two times (p<0.001) among Gypsy recruits of proven Indo-Asian origin (n=325) (in 32%) than that among Bulgarian ones (n=850) (in 16.2%). RPv incidence rate among Gypsies is of 10.4% but among Bulgarian ones is of 6%. More interesting, generalized Pv (in one third of the cases on the face, too) is by 3.6 times more common among Gypsies (in 5.29%) than

<table>
<thead>
<tr>
<th>Blood group</th>
<th>Titres of STI</th>
<th>HSV2 ≥ 1/20</th>
<th>CMV ≥ 1/20</th>
<th>RAAD</th>
<th>Rdf</th>
<th>Co.RAAD and RAAD+Rdf</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>n/%</td>
<td>n/%</td>
<td>n/%</td>
<td>n/%</td>
<td>n/%</td>
</tr>
<tr>
<td>O 157</td>
<td>21/13.4</td>
<td>26/16.6</td>
<td>30/19.1</td>
<td>0/0</td>
<td>27/17.2</td>
<td>65/41.4</td>
</tr>
<tr>
<td>A 217</td>
<td>72/33.2</td>
<td>104/47.9</td>
<td>25/11.5</td>
<td>6/2.76</td>
<td>45/20.7</td>
<td>58/26.7</td>
</tr>
<tr>
<td>B 108</td>
<td>52/48.1</td>
<td>63/58.3</td>
<td>26/24.0</td>
<td>6/5.55</td>
<td>37/34.2</td>
<td>13/13.5</td>
</tr>
<tr>
<td>AB 54</td>
<td>0/0</td>
<td>11/20.4</td>
<td>6/11.1</td>
<td>0/0</td>
<td>0/0</td>
<td>14/25.9</td>
</tr>
</tbody>
</table>

*Table 1. Relationships between ABO genetic system and sexually transmitted infections (STI) and recurrent androgen-associated dermatoses (RAAD) during navy (DN) and after navy (AN)*
among Bulgarians (in 1.44%). A similar Pv dissemination patterns are typical of the Indian population (8.9) dominated by BG-B carriers (1-3). According to the first outpatient’s attendance by students in East Africa, Pv is more common among men than among women as well as among Asian people than among individuals from other countries (10). Av sex-related tropism displays similar features (11,12).

**Blood group-related population gene pool differentiation level and its attributes: complex immune and defense (health/disease) responses**

All of these BG-associated diseases are multifactorial, i.e. a result from a complex interplay of polygenes and multiple environmental triggering factors. They are, however, not inherited in a simple Mendelian fashion and not associated with chromosomal abnormalities at all. Thus they are an area of supreme priority and challenge facing the contemporary medical genetics. Taking into consideration the basic laws of genetics formulated in 20th century (13-15) and seminal insights of Ajalla and Kriger (16), we recognize that there is another evolutionarily strong but missing key player that governs, subordinates and predetermines not only the well-known and suspected pathways of the pathogenesis and epidemiogenesis of the diseases but also the general outcome of health-disease responses’ balance. This substantial common denominator represents an epigenetically assigned major phenotype-related population gene pool (PGP) differentiation level. The latter functions either as a harmonious co-adaptive genetic (allele and interlocus) equilibrium, or as a disharmonious interlocus interaction, the so-called epistatic genetic suppression. The first one represents the normal adaptation, i.e. health by itself, and is typical of CPMs and their BG-B carriers. The second one represents the disadaptation, i.e. the pathological states or diseases by themselves are typical of TVHs as an abridged version of Western hybrid societies (WHSs). That is why WHSs turn into the most powerful generators of chronic, refractory and recurrent diseases, highly pathogenic flora carriers, infections and allergies.

The concrete PGP balance level was assessed through its sounding via some complex defensive traits such as multiple (triple or double) infectious allergic resistance (MIAR) or susceptibility (MIAS) to grippe and grippe-like conditions (GGLC), RTp, and allergy. The individual and population genetic make-up as causative factor possesses a predetermining role for human defence and immune homeostasis through the interactions between the genes. These interactions are epigenetically (evolutionarily) assigned via concrete type and intensity of the population mating as compatible inbreeding or abnormal urbanogenic interbreeding and intrinsically related to it genetic suppression. It is due to the fact that in ancient times, BG-B carriers suffered a much more severe pressure of the selection (infectious mortality) which enhanced PGP harmonization and resulted in better defense and immune capacity.

Measurements of stability and power of basic and specific cell-mediated immunity (CMI) were accomplished by intradermal testing with Candidin (C), phytohemagglutinin (PHA) and Trichophytin (T). Individuals with positive delayed skin allergy, the so-called CMI towards C and PHA at one and the same time were classified as such with co-operated basic CMI (Co.BCMI) while those with marked Co.CMI and T were considered as such with co-operative specific CMI (Co.SCMI) (Table 2). BG-B carriers who survived the plague and smallpox epidemics responded not only with a higher Co.CMI (p<0.01 - p<0.001) but also with an implied general biological compensatory reflex, i.e. with accelerated fertility. This higher fertility goes with our data showing that the parents of the examined BG-B carriers have more than two children in 75% of the cases while among those of the other blood groups this occurs only in 40-50% of the cases (p<0.001). Hence one

**Table 2. Relationships between ABO genetic system, defense polymorphism and co-operated basic cell-mediated immunity (Co.BCMI)**

<table>
<thead>
<tr>
<th>Blood group</th>
<th>n</th>
<th>Co.BCMI C+PHA n/%</th>
<th>Co.BCMI C+PHA+T n/%</th>
<th>Co.SCMI C+PHA n/%</th>
<th>Co.SCMI C+PHA+T n/%</th>
<th>MIAR triple n/%</th>
<th>MIAR double n/%</th>
<th>MIAS triple n/%</th>
<th>MIAS double n/%</th>
<th>PGP differentiation level H/D responses’ ratio %/%</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>157</td>
<td>82/52.2</td>
<td>26/16.7</td>
<td>13/8.28</td>
<td>82/52.2</td>
<td>31/19.7</td>
<td>31/19.7</td>
<td>60.5/39.5</td>
<td></td>
<td>60.5/39.5</td>
</tr>
<tr>
<td>A</td>
<td>217</td>
<td>108/49.8</td>
<td>10/4.6</td>
<td>97/44.7</td>
<td>42/19.3</td>
<td>18/8.29</td>
<td>60/27.6</td>
<td>64/36</td>
<td></td>
<td>64/36</td>
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<tr>
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<td>72/66.6</td>
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<td>66.7/33.3</td>
</tr>
<tr>
<td>AB</td>
<td>54</td>
<td>23/42.6</td>
<td>0/0</td>
<td>18/33.4</td>
<td>12/22.2</td>
<td>12/22.2</td>
<td>12/22.2</td>
<td>55.6/44.4</td>
<td></td>
<td>55.6/44.4</td>
</tr>
</tbody>
</table>
might suppose that hyperandrogeny which determined this phenomenon was coded by evolution in the genome of BG-B carriers. There is no doubt that the intense sexual potency and fertility in Eastern societies resulted in a huge population density and, logically, sufficient sex was given the status of a cult (see e.g. Camasutra and other treatises).

**CONCLUSION**

Altogether, through a 10-year longitudinal monitoring and a multitheoretical scanning of the processes (17) we establish a significant link between BG-B carriers and some STI and implied evolutionary gene-encoded hyperandrogeny and fertility, respectively. This is confirmed by a striking relationship between BG-B and long-known RAAD such as Pv and Av. We hold the opinion that ABO group polymorphism is the result from a powerful pressure of the selection by a selective BG-related infectious mortality (18-20) in ancient times. We found a confirmation of the ideas and hypotheses not only in the dramatic domination of BG-B and high sexual activity among Eastern societies but also in publishing data about a strong domination of some RAAD among them. The BG-B carriers having passed through the deadly epidemics are, logically, owners of a relatively higher Co.BCMI and Co.SCMI homeostasis and PGP-related harmonization due to a systematic tinkering and coining of evolution. Long-term prospective research is needed to better elucidate these essential topics.

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**REFERENCES**