Polymorphisms in the Promoters of MMP-1 and TIMP-1 Genes in Patients with Acne Vulgaris

H. H.Sabry\(^1\), R.M.A.Salem\(^1\), S.G.Ameen\(^2\) and S.M.Ali\(^1\)
\(^1\)Dermatology, Venereology and Andrology Dept., Faculty of Medicine, Benha Univ., Benha, Egypt
\(^2\)Clinical Pathology and Immunology Dept., Faculty of Medicine, Benha Univ., Benha, Egypt
E-Mail: Ahmed_ramadanzaid@yahoo.com

Abstract

Grid metalloproteinases (MMPs) and their particular inhibitors (tissue inhibitors of metalloproteinases, TIMPs) are fundamental for the redesigning of the extracellular framework, and incendiary infections, for example, skin inflammation. Skin break out vulgaris is an incessant incendiary illness that influences most of the populace. It is described by comedones, pustules, and papules. In this examination, it was expected to research the possible relationship between MMP-1 quality polymorphism and the hazard for skin inflammation advancement. Polymorphism of MMP-1 Gene was estimated in 60 patients with skin break out vulgaris and 40 solid controls utilizing polymerase chain response (PCR). A noteworthy connection between MMP-1 (1607 A/G) polymorphism and skin inflammation vulgaris was recognized.

Keywords: Acne vulgaris, MMP-1, Gene polymorphism.

1. Introduction

Grid metalloproteinases (MMPs) are the significant protein family liable for extracellular lattice (ECM) renovating and, all things considered, they can cut all ECM segments by sharing regular substrates. Generally, the organic job of MMPs has been connected uniquely with ECM debasement and turnover [1]. Qualities encoding collagen-debasing proteins—lattice metalloproteinase type 1 (collagenase-1) and network metalloproteinase type 3 (stromelysin)—are situated on the long arm of chromosome 11. SNP because of the inclusion of the additional guanine (G) base at position −1607/−1608 upstream from the beginning of interpretation upregulates MMP-1 translation and conceivably expands the tissue action of network metalloproteinase type 1 chemical [2].

Skin inflammation vulgaris is a constant, fiery malady of the sebaceous follicles portrayed by expanded sebum discharge and development of follicle-related shut and open comedones, papules, pustules and knobs; pseudocysts and a few sorts of scars as a sequelae of the essential sores [1].

There are four significant pathogenetic factors that have been involved in skin break out pathogenesis, in particular sebaceous organ hyperplasia with seborrhea, changed follicular development and separation, Propionibacterium acnes (P. acnes) colonization of the pilosebaceous unit, and aggravation [3].

There are four significant pathogenetic factors that have been involved in skin break out pathogenesis, in particular sebaceous organ hyperplasia with seborrhea, changed follicular development and separation, Propionibacterium acnes (P. acnes) colonization of the pilosebaceous unit, and aggravation [3].

2. Patients and methods

This is a cross Section case control study. The study included sixty patients with acne vulgaris and forty healthy subjects, age and sex matched, attending the dermatology outpatient clinic at Benha University Hospital, Benha, Egypt from May 2019 to May 2020.

Participants gave their informed written consent before enrollment and the study was approved by the Research Ethics Committee in Faculty of Medicine, Benha University. All patients included in the study have moderate to severe recurrent acne lesions, acne scar and aged from 11 to 30 years old. Patients with Acniform eruption, Acne conglobate and Fulminating acne were excluded.

2.1 Laboratory investigations

A) Two millimiters of venous blood tests pulled back from every member in the investigation into an EDTA (Ethylene Diamine Tetra-Acetic salt) tubes under complete aseptic conditions, blood was put away at - 20° until dissected.

B) DNA extraction and genotyping were done from EDTA entire blood.

DNA extraction from fringe entire blood by utilizing snappy – DNA ™ miniprep pack inventory Nos D3006, D3007, D3024 and D3025. As per maker's convention.

2.2 Statistical analysis

Measurable investigation was performed utilizing the Statistical Package for Social Sciences
Polymorphisms in the Promoters of MMP-1 and TIMP-1 Genes in Patients with Acne Vulgaris

SPSS vs.25. Mean ±SD, Tukey test, Chi square test, Fisher's careful test, Regression examination and Deviations from Hardy–Weinberg harmony were utilized.

**MMP1 (1607 A/G) alleles and genotypes in the studied groups.**

Table (1) MMP1 (1607 A/G) alleles and genotypes in the studied groups.

<table>
<thead>
<tr>
<th></th>
<th>Patients (N=60)</th>
<th>Controls (N=40)</th>
<th>P-value</th>
<th>OR</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A/A</td>
<td>N (%)</td>
<td>24 (40)</td>
<td></td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>(A/G)+(G/G)</td>
<td>N (%)</td>
<td>36 (60.0)</td>
<td>12 (30.0)</td>
<td>0.003</td>
</tr>
<tr>
<td>MMP1 (1607 A/G)</td>
<td>A</td>
<td>70 (58.3)</td>
<td>65 (81.3)</td>
<td>1</td>
<td>4.012</td>
</tr>
<tr>
<td></td>
<td>G</td>
<td>50 (41.7)</td>
<td>15 (18.8)</td>
<td>0.001</td>
<td>3.095</td>
</tr>
</tbody>
</table>

4. Discussion

Skin inflammation is a multifactorial infection that is at first determined by androgen incited expanded sebum overproduction, adjusted and strange keratinization, irritation, bacterial colonization of pilosebaceous unit on the face by Propionibacterium acnes (P. acnes), postponed type invulnerable reaction, outer components and hereditary qualities [5].

Network metalloproteinases (MMPs) are a group of proteolytic catalysts that manage extracellular grid (ECM) turnover and provocative flagging. Actuated MMPs are firmly directed by endogenous tissue inhibitor of metalloproteinases (TIMPs), which additionally apply their impacts on cell expansion, separation, apoptosis, and angiogenesis by MMP-free instruments [6].

Network metalloproteinases (MMPs) and tissue inhibitors of metalloproteinases (TIMPs) assume basic jobs in the provocative procedure and grid renovating after aggravation [7].

In this investigation, a huge connection between MMP-1 (1607 A/G) polymorphism and skin inflammation vulgaris was distinguished. The frequencies of MMP-1 (1607 A/G) genotypes and alleles were distinctive in skin break out patients and control gatherings. AG and GG genotypes and G allele appear to build the hazard to create skin break out 2-3 folds of alleles in patients and control subjects.

A similar SNP was concentrated in polycystic ovary patients. They identified a relationship between the danger of polycystic ovary and GG genotype of MMP1. Polycystic ovary is related with various skin indications identified with hormonal changes including skin inflammation vulgaris [8]. The current discoveries along with [9] results propose a hereditary foundation to this affiliation.

5. Conclusion

MMP1 (1607 A/G) quality polymorphism is related with the powerlessness of advancement of skin inflammation among the considered gathering of Egyptian populace.

II-Conflicts of interest

There are no conflicts of interest.

Reference


