1	Risk factors profile of young and older patients with Myocardial Infarction
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#### 51 Abstract

52 Myocardial infarction (MI) among young adults (< 45 years) represents a 53 considerable proportion of the total heart attack incidents. The underlying 54 pathophysiologic characteristics, atherosclerotic plaque features and risk factors profile 55 differ between young and older patients with MI. This review article discusses the main 56 differences between the younger and elderly MI patients as well as the different pathogenic mechanisms underlying the development of MI in the younger. Young 57 58 patients with MI often have eccentric atherosclerotic plaques with inflammatory 59 features but fewer lesions, and are more likely to be smokers, obese, and have poor 60 lifestyle, such as inactivity and alcohol intake. Compared to older MI patients, younger 61 are more likely to be men, have familial-combined hyperlipidemia and increased levels 62 of lipoprotein-a. In addition, MI in younger patients may be related to use of cannabis, 63 cocaine use and androgenic anabolic steroids. Genomic differences especially in the 64 pathways of coagulation and lipid metabolism have also been identified between young 65 and older patients with MI. Better understanding of the risk factors and the anatomic and pathophysiologic processes in young adults can improve MI prevention and 66 treatment strategies in this patient group. Awareness could help identify young subjects 67 68 at increased risk and guide primary prevention strategies. Additional studies focusing 69 on gene pathways related to lipid metabolism, inflammation and coagulation are 70 needed.

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72 Keywords: Myocardial infarction; Risk factors; Young patients; Atherosclerotic
73 plaque features; Genetic factors

#### 75

## 1. Introduction

Coronary artery disease (CAD) and its complications remain the most common 76 cause of death worldwide<sup>1</sup>. Evidence of elevated cardiac troponin values with at least 77 1 value above the 99th percentile upper reference limit is characterized as myocardial 78 injury<sup>2</sup>. In the 4<sup>th</sup> Universal Definition of Myocardial Infarction (MI), myocardial 79 injury differs from the term of MI. More specifically, the term MI should be used for 80 myocardial injury with clinical evidence of acute myocardial ischemia, plus the 81 detection of a rise and/or fall in cardiac troponins (cTn) values <sup>3</sup>. Additionally, one of 82 83 the following features has to be present: a) Symptoms of myocardial ischemia; b) New 84 ischemic electrocardiogram changes; c) Development of pathological Q waves; d) 85 Imaging evidence of new loss of viable myocardium or new regional wall motion 86 abnormality in a pattern consistent with an ischemic etiology; e) Identification of a 87 coronary thrombus by angiography or autopsy  $^{2,3}$ .

According to the underlying mechanisms, 5 MI Types of MI have been 88 89 recognized. Type 1 MI is presented with acute atherothrombosis in an artery which 90 irrigates a certain part of myocardium. Criteria for type 2 MI are met when an imbalance of myocardial demanded oxygen occurs <sup>4</sup>. Type 3 MI is described by cardiac death from 91 92 suspected myocardial ischemia based on electrocardiogram changes in symptomatic 93 patients, with no notification of elevated cardiac troponin levels until then. Finally, 94 Type 4 procedural MI is directly related with percutaneous coronary intervention and Type 5 with coronary artery bypass grafting  $^{4,5}$ . 95

96 CAD mainly affects older individuals as shown in the landmark Framingham 97 Heart Study, demonstrating an 8-fold increase in the MI incidence in the older age 98 group compared to participants younger than 55 years of age <sup>6</sup>. The incidence of 99 myocardial infarction (MI) among young adults has increased in the past decade as 100 reported recently in the analysis of the Atherosclerosis Risks in Communities (ARIC) study. Overall, data regarding MI incidence in younger patients are sparse, but 101 increasing numbers of studies are being conducted in this important subgroup (Table 102 103 1). Another documented difficulty is the lack of a universally established age cut-off across the published studies, making comparisons virtually impossible. For the purpose 104 of this review, an age cut-off of 45 years was used as the incidence of aggravating 105 cardiovascular risk factors such as arterial hypertension, dyslipidemia, and diabetes 106 mellitus exponentially increases past that point <sup>7, 8</sup>, accounting for excess risk factor-107 related mortality. These results highlight how challenging it is to identify risk and 108 109 genetic factors in young individuals and suggest that future research is needed to 110 identify and treat cardiovascular risk in young adults, particularly those under the age of 45 who are only rarely candidates for statin therapy <sup>9, 10</sup>. Additional studies are 111 needed to evaluate the impact of gender on clinical presentation, treatment patterns, and 112 outcomes of MI in young patients <sup>9</sup>. The mechanisms of MI in young individuals seem 113 114 to differ significantly from those affecting older patients and could be grouped into three main categories: 1) atherosclerotic coronary artery disease; 2) non-atherosclerotic 115 ischaemic heart disease; 3) hypercoagulable state. The individual's genetic profile 116 117 contributes, together with other predisposing factors, to create a favorable milieu for the development of atherosclerosis and MI at a younger age <sup>11</sup> (Figure 1). 118

119 This review article discusses the main differences that exist between the younger 120 and elderly MI patients in the four different categories mentioned above. The article 121 also addresses the different pathogenic mechanisms underlying the development of MI 122 in the younger.

123 2. Coronary plaque features – differences between elderly and younger MI
 124 patients

125 Angiographic as well as pathologoanatomical coronary findings are different in younger patients with CAD compared with the elderly (Table 2 and Figure 2). The 126 left anterior descending (LAD) is the most commonly affected artery in both patient 127 groups. Younger patients frequently present with single-vessel CAD <sup>12, 13</sup>, fewer 128 coronary lesions <sup>12</sup> and lower lesion complexity, as estimated by the Gensini score <sup>11</sup> 129 which is a strong predictor of successful restoration of myocardial perfusion <sup>14</sup> as well 130 as short and long-term adverse cardiovascular events <sup>15</sup><sup>16</sup>. Moreover, Gensini score was 131 an independent predictor of long-term mortality in elderly individuals <sup>17</sup>. 132

With regards to histological findings, younger patients were more likely to 133 have eccentric lesions, as well as lymphocytic infiltration of large- and medium-sized 134 coronary arteries and thrombosis compared with older MI patients <sup>12</sup>. Specifically, 82% 135 of the young patients had eccentric atherosclerosis with an inflammatory response being 136 observed in all of them. On the contrary, an eccentric atherosclerotic pattern was 137 detected in 39% of the elderly, while the presence of inflammation declined with aging 138 <sup>12</sup>. Plaque hemorrhage was less common in young subjects (32% vs. 61% among older 139 patients)<sup>12</sup>. Pultaceous debris, the principal component in atherosclerotic plaques of 140 141 both groups of patients, was more common in the elderly, whereas foam cells and fibrous tissue were common in both patient groups <sup>12</sup>. 142

These findings suggest that young MI patients frequently have eroded plaques <sup>18</sup>, characterized by eccentricity and infilitration by lymphocytes <sup>19</sup>. Inflammation and intraplaque hemorrhage are among the main mechanisms of vulnerable thin-cap fibroatheromas formation in the elderly, leading to plaque rupture and myocardial infarction <sup>20</sup>. Interestingly, those alterations in plaque morphology lead to distinct clinical phenotypes. An MI event in young patients is more frequently caused by plaque erosion whereas plaque rupture is more often detected in older patients <sup>21, 22</sup>. Moreover,

150 plaque erosions have been associated with a less complex, LAD-localizaed atherosclerotic pattern <sup>23</sup> as in the case of young MI patients and a better prognosis 151 compared to ruptured plaques, which are more frequently found in elderly subjects <sup>24</sup>. 152 Last but not least, the presence of plaque erosions, as in young MI patients, has 153 important therapeutic implications since a stentless, intensive antithrombotic approach 154 in a small clinical study of acute MI patients with evidence of plaque erosion led to 155 significant or complete thrombus resolution <sup>25</sup> and freedom from adverse cardiac events 156 after 4 years of follow-up <sup>26</sup>. The discovery of biomarkers of plaque erosion may 157 ultimately result in a non-invasive management of patients with non-ST elevation MI 158 159 (NSTEMI) may guide treatment decisions towards an individually-tailored treatment of intense antithrombotic regimens<sup>27</sup>. 160

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#### 3. Traditional risk factors for mi type-i in young patients

163 (The role of traditional risk factors for MI in young patients is summarized in
164 Table 3 and discussed in details below.)

*i)* Tobacco: Younger MI patients are more likely to be current smokers (80% vs. 165 57%) compared with the elderly <sup>28</sup>. Smoking is highly prevalent in ST-segment 166 elevation myocardial infarction (MI) in young patients <sup>29</sup>. A dose-effect between 167 smoking and MI is present with patients who smoked >25 cigarettes/day having 8-fold 168 higher odds of MI compared to never smokers <sup>30</sup>. Both cigarette and waterpipe smokers 169 170 were more common among younger first-time MI patients than older first-time MI patients <sup>31</sup>. Smokeless tobacco has been associated with lower high-density lipoprotein 171 (HDL) and higher total cholesterol levels and can potentiate coronary vasoconstriction, 172 therefore possessing atherogenic and thrombogenic properties <sup>32</sup>. The detailed 173 mechanism through which cigarette smoking is associated with cardiovascular disease 174

has not yet been clarified. It is strongly suggested that smoking has two effects on
platelets: a) a significant acute potentiation of platelet activation occurring shortly after
smoking a cigarette b) a chronic desensitization of the cell to activating agents occurring
during the period between cigarettes leading to Type 1 MI <sup>33, 34</sup>.

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180 ii) Dyslipidemia: Dyslipidemia is an established MI risk factor among all age groups. Hovingh et al. reported a high prevalence (10%) of familial-combined 181 hyperlipidemia (FCHL) in survivors of premature MI, while the levels of low-density 182 lipoprotein (LDL-C) remained >70mg/dl irrespective of statin use <sup>35</sup>. More specifically, 183 FCHL was associated with a 24-fold increased adjusted risk for MI with very-low-184 185 density lipoproteins and non-high-density lipoprotein constituting aggravating factors for MI incidence <sup>36</sup>. FCHL is often present in patients with a family history of premature 186 CAD (20% prevalence in young MIs) and/or high levels of LDL-C (60% prevalence 187 "young" MIs)<sup>36</sup>. Dyslipidemia has a stronger correlation with Type 1 MI incidence in 188 189 the elderly compared with younger individuals (43% vs 36%). However, crosssectional studies found that triglyceride, LDL-C and apolipoprotein B levels were 190 significantly higher in younger compared with older MI patients, whereas HDL-C 191 192 levels were lower <sup>37</sup>.

Adolescents with a parental history of premature myocardial infarction have increased lipoprotein-a (Lp(a)) levels. Likewise, a high level of Lp(a) has described as an independent risk factor for MI in all age groups <sup>38</sup>. Rallidis et al. showed that high levels of Lp(a) increase by 3-fold the odds of MI in individuals <45 years, with a lesser association in individuals between 45 and 60 years. An increase of 10mg/dl results was associated with a 4% higher relative risk of having Type 1 MI at a younger age (<45 years) and 2% in middle age (45-60 years) <sup>39</sup>. 200

201 *iii) Obesity:* Liu et al. analyzed the correlation between Healthy lifestyle factors (HLFs) and MI in younger patients, such as 1) average body mass index <25kg/m<sup>2</sup>; 2) 202 No or moderate alcohol intake; 3) higher healthy diet score; 4) higher physical activity 203 204 score, and 5) Never smoking. The prevalence of CAD in combination with age, sex and 205 race were 3.0%, 14.6%, 29.5%, 39.2% and 60.7% for people with 0-1, 2, 3, 4, and 5 206 HLFs respectively with similar graded relationships being observed for each sex-race group <sup>40</sup>. Compared with older (>45 years) MI patients, younger patients were more 207 likely to be male and had higher body mass index  $(31 \text{kg/m}^2 \text{ vs. } 29 \text{kg/m}^2)^{40}$ . 208 209 Adiposopathy is comprised by adipocyte hypertrophy, decreased adipose tissue blood 210 flow, altered oxygen levels within the tissue, a state of chronic low-grade inflammation and blunted lipid metabolism <sup>41,42</sup>. With obesity levels increasing, the risk of premature 211 Type 1 MI is likely continue increasing, with further research needed on the effect of 212 adiposopathy on young subjects <sup>43</sup>. 213

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iv) Sex: Most young MI patients are men. Young men have higher levels of 215 cardiac biomarkers and more classic electrocardiogram findings while women tend to 216 present with more atypical symptoms and fewer STEMI cases <sup>44, 45</sup>. Specifically, among 217 218 patients with MI presenting with chest pain, female patients reported more often 219 additional symptoms, such as palpitations, shortness of breath, and epigastric pain. 220 Moreover, females often interprete their anginal symptoms as high anxiety levels and 221 this results in delayed hospital presentation and potentially worse prognosis of female patients with acute MI<sup>44</sup>. 222

Recent data suggest a rising incidence of MI among young women withspontaneous coronary artery dissection being the cause in a sigficant proportion of

STEMI <sup>45, 46</sup>. Young women with MI are more likely to have chronic obstructive pulmonary disease, congestive heart failure, morbid obesity, diabetes, hypertension and renal failure while they usually suffer from higher levels of anxiety and have altered mental health and physical status <sup>47</sup>. As far as changes in trends are concerned, the prevalence of hypertension has increased while smoking was less frequently observed in young women in the course of time <sup>45</sup>.

Women experience their first MI 6-10 years later than men and a protective effect of their natural estrogen status prior to menopause has been suggested. Female sex hormones have been associated with a less atherogenic lipid profile and a more healthy fat distribution <sup>48</sup>. Several studies are trying to identify the protective role of estrogens on cardiovascular system with further research needed <sup>49</sup>.

With regards to management, important disparities are present between sexes. Specifically, young women are less likely to undergo an early invasive strategy, primary percutaneous coronary intervention or CABG compared to their male counterparts irrespective of MI type, even though improvements are noted in the course of time <sup>50</sup>. This translates into worse prognosis (in-hospital mortality, vascular complications, major bleeding) compared to male patients. Interestingly, increasing age in females was associated with improved outcomes in comparison to elderly males <sup>50</sup>.

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244 *v) Diabetes mellitus:* Although diabetes mellitus (DM) is rare in young patients 245 with MI, it is nevertheless associated with a higher MI risk in both sexes <sup>47</sup>. DM, 246 hypertension, dyslipidemia and the previous history of MI were more common among 247 elderly patients, 37%, 60%, 43%, and 42%, respectively, versus 10%, 24%, 36% and 248 25% in the younger population <sup>28, 51</sup>. In the setting of type 1 DM, age of onset and 249 gender are important determinants of survival and MI outcomes in young subjects.

250 Specifically, women who developed type 1 DM before 10 years of age had a hazard 251 ratio of 91.07 with the corresponding hazard ratio in men being 15.11. These differences, however, were mitigated with increasing age of onset, as the lowest hazard 252 253 ratio for MI in women with type 1 diabetes was observed with disease onset between 26-30 years <sup>52, 53</sup>. The above-mentioned finding could be explained by the action of 254 255 glycosylated hemoglobin A1c, which is described as an independent determinant for microvascular perfusion, suggesting tight glycemic control is potentially important for 256 the prevention of cardiovascular disease <sup>54-56</sup>. 257

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# a) Anabolics and stimulants

4. Risk factors for MI at a younger age

261 Substance abuse, such as cocaine and cannabis, is among the less common MI risk factors for MI<sup>57</sup>. Specifically for cannabis, which is the most frequently abused 262 263 substance, it has been linked with incident MI independently of traditional 264 cardiovascular risk factors, with the effect being more pronounced in younger patients <sup>58</sup>. Similarly for cocaine use, which is reported in approximately 10% of MIs at a young 265 age, has been associated with an increased risk of cardiovascular mortality. The 266 267 responsible mechanisms include increased myocardial oxygen demand, lower peripheral vascular reflex response, as well as coronary artery vasospasm, meeting the 268 Type 2 MI criteria. The risk of MI due to cocaine is dose-independent, in contrast to 269 cannabis 58. 270

Moreover, the use of stimulating substances, mainly androgenic anabolic steroids (AASs) among elite as well as amateur athletes is widespread, with high doses leading to numerous side-effects. AASs increase the odds for MI in young patients due to a decrease in HDL-C and Apolipoprotein A1 (by 20-70%) and a significant increase of

LDL-C and apolipoprotein B (up to 20%) 59. Besides, long-term use of AASs could 275 276 lead to the development of hypertension and a high concentration of C-reactive protein 277  $^{59}$ . The main pathway of thrombosis is due to erythrocytosis (9.6% increase in 278 hematocrit within approximately 26 weeks of use), thrombocytosis and platelet hyperactivity <sup>59</sup>. Furthermore, AAS increase levels of procoagulant factors (especially 279 280 fibringen, factor VIII and X), homocysteine as well as endothelium release of proteins C and S, with the decreased fibrinolytic activity (decreased levels of a-2-macroglobulin 281 282 and plasminogen activator inhibitor 1 as well as increased levels of tPA and 283 plasminogen) and prostacyclin synthesis further enhancing their prothrombotic action 59. 284

285 Erythropoiesis-stimulating agents (ESAs) are erythropoietin derivatives have been widely used as performance-enhancing drugs. Induced erythrocytosis can be 286 achieved through the use of erythropoietin and analogs, blood transfusion in the form 287 288 of homologous or autologous administration as well as by Red Blood Cells-mimicking synthetic biomaterial particles <sup>60</sup>. There is limited data regarding ESAs in the setting of 289 290 CAD and MI, but it is speculated that their combination with dehydration during physical activities could lead to adverse cardiovascular events <sup>60</sup>. According to the 291 292 above mentioned, using of AASs and ESAs lead to acute coronary atherothrombosis via multiple pathways, following the Type 1 MI criteria<sup>61</sup>. 293

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#### b) Thrombotic / Fibrinolytic factors

Thrombotic and fibrinolytic pathways are complex and display a particular interaction between them. Redondo et al. showed that high levels of factor V or factor VII in serum plasma are associated with higher MI risk. The presence of smoking or arterial hypertension magnified the risk by 50-fold <sup>62</sup>. Factor V Leiden, the most

300 common hereditary hypercoagulability factor has been associated with premature MI Type 1, while a higher activity of factor XIII was also detected in young MI survivors. 301 Moreover, Factor XI can activate coagulation factors X, V and VIII, and inhibit the 302 303 anti-coagulant tissue factor pathway inhibitor, therefore being recognized as an independent risk factor for MI<sup>63</sup>. The higher risk associated with these hypercoagulable 304 305 states could be related to unfavorable lipidemic profile, as demonstrated by abnormal Lp(a) levels in a group of patients with antiphospholipid syndrome <sup>62, 64, 65</sup>. 306 307 Furthermore, Butt et al. demonstrated that the Factor II 20210A allele, the Factor XIII-A Leu34 allele and their synergistic effect are additional risk factors for MI <sup>62, 66</sup>. A 308 recent meta-analysis demonstrated an unfavorable role of hypercoagulable states with 309 310 previous MI without such an association in the setting of stable CAD <sup>67</sup>.

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312 c) Homocysteine

313 Hyperhomocysteinemia causes the production of proinflammatory cytokines, 314 namely interleukin-1 $\beta$  and -6, tumor necrosis factor- $\alpha$ , monocyte chemoattractant protein 1 and intracellular adhesion molecule-1, leading to increased oxidative damage 315 <sup>68</sup>. There are conflicting reports on the association between homocysteine -or its 316 317 lowering with treatment- and the incidence of CAD. Numerous studies showed that hyperhomocysteinemia is associated with increased MI risk, classifying it as an 318 independent risk factor and a possible marker of preclinical disease state <sup>68, 69</sup>. Some 319 320 studies have shown higher homocysteine levels in younger infarcted patients compared with the elderly  $^{51, 70}$ . 321

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#### d) Genetic factors

324 There is strong evidence that CAD in early life is associated with the patient's genetic background. The non-Mendelian heritability of MI and CAD makes the issue 325 more complex. A study in 2017 -which examined prothrombotic risk factors- showed 326 327 that polymorphisms G20210A of prothrombin [FII] gene are associated with increased risk of premature ST-segment elevation MI<sup>71</sup>. Prothrombin (FII), the precursor of 328 thrombin, is a vitamin K-dependent glycoprotein whose primary function is to convert 329 fibrinogen to fibrin, activating factor XIII in the development of clots that are more 330 resistant to fibrinolysis. The expression of the mutation G20210A results in slightly 331 higher levels of prothrombin, which can be easily converted to thrombin as required, 332 predisposing to hypercoaguable state <sup>72</sup>. If the above polymorphism is combined with 333 casual smoking, the risk is increased 22-fold (95% CI: 9.192-66.517)<sup>71</sup>. Hmimech et 334 al. showed that polymorphisms G20210A of prothrombin [FII] gene -even if there is a 335 single or double copy of the 20210A allele- is highly associated with premature MI<sup>73</sup>. 336 337 A recent meta-analysis found that the polymorphism raises MI risk in an age-related way, with youngs under the age of 55 experiencing the most (OR = 1.76, 95%CI: 1.32-338 2.35) <sup>74</sup>. Burzotta et al.'s meta-analysis showed that the G20210A prothrombin gene 339 340 polymorphism can be a minor but substantial risk factor for myocardial infarction at a 341 young age (< 45 years) (OR=2.3, CI: 1.27–4.59), favoring the expression of ischaemic 342 cardiac disease in persons with a small degree of coronary atherosclerosis on angiography <sup>75</sup>. Other polymorphisms that seem to play a significant role are these of 343 factor V Leiden - especially the homozygote phenotype-, plasminogen activator 344 345 inhibitor 1 polymorphism 4G/5G and glycoprotein VI (GP6, 13254 TC, Ser219Pro). Generally, FV activates factor X in the process of transforming prothrombin to 346 347 thrombin. FV is activated by thrombin, and its active form is vulnerable to protein C 348 cleavage and inactivation. The V Leiden mutation altered FV structure, leading to

protein C tolerance with a longer half-life and increased thrombin production <sup>76</sup>. 349 350 Mannucci et al. showed that the minor A allele of F5 G1691A was associated with an increased risk of MI, noting the important role of hypercoaguability in the pathogenesis 351 of MI in young individuals (<45 years) <sup>77</sup>. Studies in youngs under 45 years old with V 352 353 Leiden mutation showed that the risk for MI was 32-fold higher in smoking than the non-smokers, but no sex association has been proved until now <sup>78, 79</sup>. Congenital 354 355 deficiencies of AT, PC, and PS belong to rare diseases, the few available case reports refer to young individuals with MI. This could imply that hereditary deficiencies of 356 357 the natural anticoagulants are significant enough to induce early onset of thrombotic complications<sup>80</sup>. The above mentioned suggests that thrombophilia might be an 358 359 important part of the differential diagnosis in MI cases with otherwise unexpected 360 coronary disease occurring at a young age.

Titov et al. analyzed genotype frequencies of single nucleotide polymorphisms 361 362 (SNPs) in genes whose protein products are involved in the pathogenesis of 363 atherosclerosis. In the group of age <60 years SNPs of FGB, TGFB1, ENOS and CRP are associated with high risk of MI, with the higher risk to be observed in their 364 combinations [FGB + TGFB1; FGB + LPL + IL4; FGB + ENOS]<sup>81</sup>. The genome-wide 365 association studies (GWAS) identified 6 new loci associated with CAD: on 2q37 366 (KCNJ13-GIGYF2), 6p21 (C2), 11p15 (MRVI1-CTR9), 12q13 (LRP1), 12q24 367 (SCARB1), and 16q13 (CETP) with odds ratio per copy of the risk allele ranging from 368 1.04 to 1.09<sup>82</sup>. The ADAMTS7 polymorphism exhibited a significant effect on MI risk, 369 370 with hypertensives, non-diabetics and patients with hyperlipidemia possessing the greatest risk <sup>82</sup>. Analysis of 2,967 early-onset MI cases identified SNPs at nine loci: 371 372 three are newly identified (21q22 near MRPS6-SLC5A3-KCNE2, 6p24 in PHACTR1 and 2q33 in WDR12) and six replicated prior observations <sup>83-85</sup> (9p21, 1p13 near 373

374 CELSR2-PSRC1-SORT1, 10q11 near CXCL12, 1q41 in MIA3, 19p13 near LDLR and 375 1p32 near PCSK9<sup>86</sup>. The role of ADAMTS-7 in neointima formation is mediated via increased smooth muscle cell (SMC) migration caused by cartilage oligomeric protein 376 377 degradation and impaired re-endothelialization. Higher levels of ADAMTS-7 correlated with high levels of lipid content, but with low smooth muscle cell and 378 379 collagen content in atherosclerotic plaque formation, both of which are indicators of a vulnerable phenotype <sup>87</sup>. Further studies are needed to clarify the exact mechanism for 380 381 this association.

382 As discussed above, high Lp(a) levels are associated with incident MI. Lp(a) prevents endothelial cell plasmin production and disrupts the fragile equilibrium 383 384 between thrombus formation and fibrinolysis. Lp(a)'s gene is one of the strongest 385 monogenic risk factors for CAD. Clarke et al. identified two Lp(a) variants associated with increased levels, located on chromosomal region 6q26-27 with odds ratio for 386 coronary disease being 1.70 and 1.92, respectively <sup>88</sup>. Genetic data support Lp(a) levels 387 388 < 20 mg/dl as optimal, and as atherothrombotic range > 30-50 mg/dl. Clarke and 389 pioneers established the causative relation between Lp(a)'s genotype and CAD via extended genetic research <sup>88-91</sup>. Methylenetetrahydrofolate reductase polymorphism-390 391 homozygotes (C677T MTHFR and 1298 AC) with or without casual smoking is a major 392 risk factor, whereas heterozygotes have no statistically significant association with 393 premature MI. According to the findings of a recent meta-analysis, the MTHFR C677T 394 polymorphism is associated with an increased incidence of MI in young/middle-aged Caucasians. This connection could not be established in the elderly population <sup>92</sup>. In the 395 group of people with a mutation in the gene MTHFR with alleles C677T or A1298C -396 397 particularly the homozygotes- has been observed an increase of their plasma 398 homocysteine and the vascular damage, but exceptions have been published for these

399 mutations, with absence of elevated homocysteine levels in countries where food is 400 fortified with folic acid, i.e. the United States <sup>93-95</sup>. The European Prospective Cohort 401 on Thrombophilia findings indicated no substantial elevated risk of mortality in 402 individuals neither those with thrombosis occurrence <sup>96</sup> (**Table 4**) (**Central** 403 **Illustration**).

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#### 5. Clinical Implications

Taking into account the differences in the profile of young and older MI patients, 406 customization of the established primary and secondary prevention strategies may be 407 408 considered. Since, it is rather difficult to identify hereditary atherosclerotic burden in 409 young population, clinicians have to prevent the thrombotic events. Clinical awareness 410 for the development of CAD even at younger age is required. The value of 411 asymptomatic screening for CAD has to be explored especially in subgroups of patients 412 with strong family history of premature CAD or presence of non-traditional risk factors 413 (e.g. increased lipoprotein(a) or C-reactive protein serum levels). Surveillance and monitoring for the early onset of arterial hypertension, an abnormal lipid profile, central 414 obesity and strong counselling against smoking should be implemented, especially 415 416 when other predisposing thrombotic factors co-exist e.g. use of oral contraceptives in young women <sup>97</sup>. Clinicians must be conscious of the prevalence of drug abuse among 417 young people, which has been steadily growing over the years <sup>98</sup>. According to studies, 418 419 the majority of young people with premature CAD would not have received statin medication prior to their first MI event 99-101. This would be disastrous, evidence 420 suggests that the advantage of statin therapy increases with treatment duration; young 421 422 people with early onset of CAD may derive the largest benefit from timely 423 administration of preventive measures (like statins) given the higher average life

424 expectancy <sup>99-101</sup>. The declining costs of genetic tests and whole-genome sequencing
425 may also lead to the implementation of genetic screening to patients with strong family
426 history of premature CAD.

427 Younger people are also less likely to be recommended for statin therapy following a MI injury than older people, despite having a significantly greater potential 428 life expectancy for repeated events <sup>102</sup>. These findings point to the need for active 429 secondary preventive interventions in young adults who suffer from MI, especially 430 young patients, who are only rarely candidates for statin therapy. Even so, no clinical 431 432 trials have been conducted to evaluate the positive benefits of anticoagulation in patients with hereditary thrombophilia and arterial thrombotic events such as MI<sup>103</sup>. A 433 434 timely treated incident has better outcome and prognosis for young counterpants <sup>104</sup>. 435 The major area of concern is that approximately 50% of total young MI patients may not receive any reperfusion therapy due to late diagnosis <sup>105</sup>. Due to a more patent 436 437 infarct-related artery and non-significant disease in non-infarct vessels, young MI 438 patients have a considerably greater likelihood of revascularization when thrombolysis occurs within 6 hours of an event <sup>105</sup>. On the other hand, developments in plaque 439 imaging provide the opportunity to discern the intrinsic pathophysiological cause of 440 thrombosis <sup>106</sup>. Plaque erosion, the dominant mechanism of coronary thrombosis in 441 young individulas, can now be reliably detected by intracoronary imaging, implying 442 that treatment could be individualized based on pathophysiology of MI; 443 pharmacological rather than mechanical intervention could provide an optimal 444 treatment for patients with plaque erosion <sup>106, 107</sup>. The EROSION study showed that MI 445 patients with eroded plaques receiving antithrombotic therapy without stenting (heparin 446 447 for 3 days with concurrent aspirin and ticagrelor, glycoprotein IIb/IIIa antagonists, tirofiban) showed > 50% decrease in thrombus volume and approximately no major 448

adverse cardiac events occurred <sup>25</sup>. Larger, randomized studies will confirm this propotition and evaluate the use of antithrombotic treatment in cases of eroded plaques <sup>107</sup>. On the other hand, a randomized trial in 2020 showed that prophylactic stenting of vulnerable plaques (lesions with visually-estimated diameter stenosis 40%, but with plaque burden by intravascular ultrasound of  $\geq$ 65%) was safe with enlarged minimum lumen area and favorable clinical outcomes (MACE) during the follow up <sup>108</sup>.

455

456 **6.** Conclusion

457 Young MI patients have a cluster of risk factors including eccentric atherosclerotic plaques with inflammatory features, higher incidence of tobacco use, 458 obesity and increased healthy lifestyle risk factors such as inactivity and alcohol intake. 459 460 Compared with older patients where MI is prevalent among men and women, young MI patients are more likely to be men, have a family history of familial-combined 461 462 hyperlipidemia and higher levels of lipoprotein(a). In addition, cannabis and cocaine 463 use, as well as the use of androgenic anabolic steroids are risk factors for MI in young patients. Genomic differences, especially in the pathways of coagulation and lipid 464 465 metabolism, have also been identified between young and older patients with MI. The 466 relative contributions of gene pathways related to lipid metabolism, inflammation, 467 cellular proliferation, vascular tone, or other as yet undiscovered pathways may provide 468 important insights. Both familial hypercholesterolemia mutations and high polygenic 469 scores are associated with increased odds of early-onset MI. The different pathophysiology and risk factor profile of young and older MI patients could help 470 471 identify young subjects at increased risk and guide primary and secondary prevention 472 strategies.

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- 480 None
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992 Legends

993 Figure 1: Pathophysiology of myocardial infarction among young adults.

- 994 MI: Myocardial Infarction; CRP: C-Reactive Protein; PAPP-A: Pregnancy associated
- 995 plasma protein-A; sCD40L=Soluble CD40 ligand; oxLDL: oxidized Low Density
- 996 Lipoprotein; MPO: Myeloperoxidase; MCP-1: Monocyte Chemoattractant Protein-1;
- 997 MMPs: Matrix Metalloproteinases; sICAM: Circulating Soluble Intercellular Adhesion
- 998 Molecule; sVCAM: Circulating Soluble Vascular Cell Adhesion Molecule; IL-1/IL-
- 999 6/IL-18: Interleukin-1/-6/-18; TNF-a: Tumor necrosis factor-a; LA: Lupus
- 1000 anticoagulant; aCL: Anticardiolipin antibodies; Anti-b2–GPI: Anti-beta2-glycoprotein
- 1001 I; Fc gamma RIIA: Fc gamma RIIA/CD32a Recombinant Protein Antigen; Anti-PF4:
- anti-platelet factor 4
- 1003 Figure 2: Comparison of angiographic features between young and elderly1004 patients.
- 1005 IL-1/IL-6/IL-18: Interleukin-1/-6/-18; TNF-a: Tumor necrosis factor-a; MCP-1:
- 1006 Monocyte Chemoattractant Protein-1
- 1007 Central Illustration: Differences in risk factors profiles between young and elderly
- 1008 patients with MI
- 1009 Arrows illustrate frequency of occurrence of each risk factor in each age goup.
- 1010

# Table 1. Comparative incidence of MI in young and old patients according to completed studies

Study	Country	Time interval	Young age cut-off	Young MI, %	Old MI, %
ARIC <sup>45</sup>	USA	1995-1999		27	73
		2000-2004	<55 years	30	70
		2005-2009		32	68
		2010-2014		32	68
Qureshi et al <sup>109</sup>	USA	2005-2014	<55 years	30	70
Odoi et al <sup>110</sup>	USA	2005-2014	<45 years	4.3	95.7
Wittlinger et al <sup>111</sup>	Germany	2005-2014	<40 years	2	98
	New Zealand	2006	<45 years	3.3	96.7
Wang et al <sup>112</sup>		2016		2.9	97.1
Jortveit et al <sup>113</sup>	Norway	2013-2016	<45 years	4.4	95.6
Schmidt et al <sup>114</sup>	Denmark	1989-1998	<50 years	7.5	92.5
		1999-2008		8	92
Seo et al <sup>115</sup>	Korea	2006-2010	.50	15.6	84.4
		2011-2015	<50 years	14	86
Alkhouli et al <sup>50</sup>	USA	2003-2015	<45 years	10.5	89.5

CVR-2020-1966-R1 MI=myocardial infarction, USA=United States of America

# Table 2: Coronary plaque features of young and older MI patients.

	Plaque feature	< 45 years	> 45 years	Clinical Implications
S C		No disease	Two	
ANGIOGRAPHIC FINDINGS	Number of infarcted arteries	Or	Or	
		Single vessel <sup>13</sup>	More vessels <sup>13</sup>	Angiographic complexity and severity of coronary artery disease is
	Number of lesions	2 or less $^{12}$	3 or more $^{12}$	associated with increased in-hospital, short- and long-term MACE
	Gensini score	$7.69 \pm 5.23$ <sup>11</sup>	$16.08 \pm 7.81$ <sup>11</sup>	15, 116-118
	Plaque eccentricity	82% 12	<b>39%</b> <sup>12</sup>	
TICS	Inflammatory response	100% 12	61% <sup>12</sup>	
PLAQUE CHARACTERISTICS	Thrombosis	47% <sup>12</sup>	11% 12	Young MI patients frequently present with plaque erosions (eccentric with lymphocytic inflammatory infiltrate), which are met
	Plaque hemorrhage	32% 12	61% <sup>12</sup>	with a favourable prognosis and an opportunity for conservative
	Principal component	39% pultaceous debris 25% foam cells <sup>12</sup>	64% pultaceous debris	management with intense antithrombotic treatment.
			18% dense fibrous	
PL			tissue <sup>12</sup>	

MACE=major adverse cardiovascular events, MI=myocardial infarction

Die 5. Differences a	nd similarities in risk f			patients with with
<b>Risk Factor</b>	Studies	Age <45 years	Age >45 years	Results
Smoking	Shah et al. <sup>119</sup> Hbejan <sup>30</sup> Larsen et al. <sup>29</sup> Deligiannis et al. <sup>32</sup> Liu et al. <sup>40</sup> Zgheib et al. <sup>28</sup>	(+)(+)(+)(+)	(+)(+)	A dose-effect response is associated with young MI incidence. Younger patients with MI were more likely to b smokers (80% vs. 57%) compared to the elderly.
Male gender	Bucholz et al. <sup>47</sup> Shah et al <sup>119</sup> Liu et al. <sup>40</sup>	(+)(+)(+)(+)	(+)(+)(+)	The patients of the young MI group were more likely to be male (80%). The dominance o men versus women is frequently reported (80% vs 71%).
Diabetes mellitus	Zgheib et al. <sup>28, 51</sup>	(+)	(+)(+)	DM is more common among elderly patients (10% vs 37%)
Obesity (BMI>25kg/m <sup>2</sup> )	Matsis et al. <sup>120</sup> Liu et al. <sup>40</sup> Shah et al. <sup>119</sup>	(+)(+)(+)(+)		80% of young patients with M have a higher BMI. Compared to older MI age group (>45 years), patients in the younger group were more likely to be male and have a higher BMI (31kg/m <sup>2</sup> vs. 29kg/m <sup>2</sup> ).
Homocysteine	Karim et al. <sup>51, 70</sup>			Homocysteine levels are increased in younger infarcted patients compared to the elder $(16.36 \pm 7.8 \text{ mmol/l vs. } 11.7 \pm 5.6 \text{ mmol/l}).$
Hypertension	Zgheib et al. <sup>28, 51</sup>	(+)	(+)(+)(+)	Hypertension is more common among elderly patients (24% v 60%).
Dyslipidemia	Zhang et al. <sup>37</sup>	(+)(+)	(+)(+)	Dyslipidemia has a stronger correlation with MI incidence the elderly compared to young individuals (43% vs 36%).
FCHL	Hovingh et al. <sup>35</sup> Singh et al. <sup>121</sup> Shah et al. <sup>119</sup> Wiesbauer et al. <sup>36</sup>	(+)(+)	(+)	FCHL reduced the age onset of first MI by as much as 15 year Approximately 10% of young MI patients present with elevated LDL-C levels. The EUROASPIRE IV cohort stud of 7044 patients with MI, showed that 8.3% had probabl FCHL, increasing to >15% in patients with premature event.
PHPMI	Rallidis, L.S., et al $_{39}$ Gaeta, G., et al. $^{38}$ Shah et al. $^{119}$	(+)(+)(+)		Lp(a) levels are increased in healthy young patients with parental history of premature MI. PHPMI is reported in the

				vast majority of young MI cases (41%-71%).
Previous MI	Zgheib et al. <sup>28, 51</sup>	(+)	(+)(+)	Incidence of previous MI is more common in the elderly (25% vs 42%).
Hereditary Coagulopathies/ Genetic mutations	Redondo et al. <sup>62, 63</sup> Clarke et al. <sup>80</sup>			Latest studies demonstrate that hereditary coagulopathies have a significant association with premature MI. Genetic mutations are part of differential diagnosis in cases with unexpected CAD occurring at a young age.
Illicit/performance- enhancing drugs	Lisowska et al <sup>57</sup> Deligiannis et al. <sup>32</sup>			Cardiovascular toxicity includes atherogenic, thrombotic and hematological effects as well as direct myocardial injury.
<u>Lp(a)=lipoprotein-a, =</u> 1013 1014 1015	•			ellitus, BMI=body mass index,
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Gene of	Polymorphism- Mutation	plicated in MI incidence. Studies report on MI risk in Young population	OR	Action
Factor II	G20210A <sup>73</sup>	<ul> <li>Li et al, 2017<sup>74</sup></li> <li>Burzotta et al, 2004<sup>75</sup></li> </ul>	<ul> <li>Particularly in youngs (≤55 y) and in Caucasians (OR= 1.76, 95% CI: 1.32–2.35)</li> <li>Moderate risk factor in young individuals (≤45 y) (OR=2.3, CI: 1.27–4.59)</li> </ul>	Hypercoagulation
Factor V	V Leiden (rs6025) <sup>80</sup>	<ul> <li>Juul et al, 2002<sup>122</sup></li> <li>Mannucci et al, 2010<sup>77</sup></li> </ul>	<ul> <li>Potential moderate risk factor in young individuals (&lt; 50 y) (OR=1.54, 95% CI: 1.07–2.22)</li> <li>Moderate risk factor (&lt;45y) (OR=1.61, 95% CI: 1.16–2.22)</li> </ul>	Hypercoagulation
Antithrombi n Deficiency	p.Arg79Cys p.Pro73Leu nt9788G>A g.5924delC	<ul> <li>Several Cases Reports</li> </ul>	NA	Hypercoagulation
Protein C Deficiency	p.Pro369Leu p.Gly109Arg p.Lys193del c.565C/T p.Arg147Trp	<ul> <li>Several Cases Reports</li> </ul>	NA	Hypercoagulation
Protein S Deficiency	p.Thr673fsX10 p.Ser501Pro (Heerlen) p.Asp496	<ul> <li>Several Cases Reports</li> </ul>	NA	Hypercoagulation

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Methylenete trahydrofola te reductase	C677T MTHFR or 1298 AC <sup>93-95</sup>	• Chao et al, 2011 <sup>92</sup>	<ul> <li>Moderate risk in young/middle- aged (&lt;50 y) Caucasians (OR = 1.275, 95% CI: 1.077- 1.509)</li> </ul>	Increase of plasma homocysteine.
Lp(a)	rs10455872 or rs3798220 <sup>88</sup>	NA	NA	Lp(a) variants
ADAMTS7 tagSNP	rs3825807 <sup>82</sup>	NA	NA	Affects ADAMTS7 <sup>emon</sup> maturation.
FGB TGFB1 ENOS CRP	rs1800788*T <sup>81</sup> rs1982073*T/T <sup>81</sup> rs2070744*C <sup>81</sup> rs1130864*T/T <sup>81</sup>	NA	NA	maturation. Their protein products are involved in the pathogenesis of atherosclerosis. Their products are involved in the pathogenesis of atherosclerosis.
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