Could cardiovascular disease risk stratification and management in rheumatoid arthritis be enhanced?

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The markedly enhanced risk of atherosclerotic cardiovascular disease (CVD) in rheumatoid arthritis (RA) is well documented.1 2 This prompted a European League Against Rheumatism (EULAR) task force to make a commendable effort in producing recommendations for cardiovascular risk management in patients with inflammatory arthritis.3 These included the application of the systematic coronary risk evaluation score (SCORE), a multiple major traditional risk factor assessment equation. In addition, the EULAR task force recommended applying a multiplier of 1.5 in patients with RA that met 2 of 3 criteria consisting of (1) a disease duration >10 years, (2) rheumatoid factor or anticyclic citrullinated peptide positivity and (3) the presence of extra-articular manifestation, thereby creating the modified (m)SCORE.

Risk factor assessment algorithms, including the SCORE and the Framingham risk equation, are recommended worldwide as part of CVD risk management in the population at large.4 5 These equations allow for stratifying subjects into low, intermediate, high and very high risk groups. With regard to CVD risk management, lifestyle factors should be addressed in all individuals. The use of cardiovascular drugs, particularly antihypertensive and lipid-lowering agents should be considered in those at high or very high risk as these interventions markedly reduce CVD event rates in such persons. Patients with established CVD, diabetes and chronic kidney disease are at high or very high risk and hence, risk factor equation application is not indicated.

Nonetheless, approximately a third of CVD events are not attributable to major CVD risk factors.6 Congruent with this, although multiple risk factor equations are useful in determining the overall CVD risk among different populations, they often underestimate the actual risk in individual subjects. This is particularly evident in those who are at moderate risk according to major risk factor assessment equations.4 5 Consequently, based on reported evidence, two approaches that can refine CVD risk stratification are currently considered helpful in both European and American guidelines on CVD risk management.6 7 First, the use of biomarkers, particularly high-sensitivity C reactive protein concentrations, and second, cardiovascular imaging, including multidetector tomography coronary artery calcification scores (CACS) and carotid ultrasound; for the first time, carotid plaques are recognised to represent very high risk in the latest European guidelines on CVD prevention in clinical practice,4 where cardiovascular risk assessment is also not necessary because the patient is categorised into secondary prevention equivalent to patients with established CVD. Taking into account that patients with RA have two to three times more frequent asymptomatic carotid atherosclerosis compared with persons without RA,7-12 this will have a major impact on correct classification and hence treatment in patients with RA.

Would traditional risk factors as included in the mSCORE be expected to reliably reflect the actual CVD risk in RA? Recently reported evidence suggests that this may not be the case. One study showed that the mean Framingham score was as low as 7% (low risk) in RA patients with carotid artery plaque despite it being associated with a 10-year incident CVD event rate of ∼39%.10 Indeed, Crowson and colleagues11 recently reported that the risk of CVD events is twofold and 65% higher than the Framingham score predicts in women and men with RA, respectively. The Reynolds risk score that additionally includes high-sensitivity C reactive protein concentrations and a family history of premature CVD revealed similar shortcomings.11 Using the area under the receiver operating characteristic curve in logistic regression analysis, traditional and non-traditional CVD risk factors associated similarly and additively with plaque prevalence.14 Solomon and colleagues subsequently confirmed that the same applied when the risk of incident CVD events was examined. The associations of traditional risk factors with CVD event rates are weakened in RA.16 Indeed, the relation between lipids and CVD is at most inconsistent17 and in one report was in fact paradoxically inverse, in RA.18 A recent investigation disclosed that in contrast to their white counterparts, black Africans with RA experience no major conventional risk factor–atherosclerosis and systemic inflammation–atherosclerosis relations.19 This further argues against the reliance on risk factor profiles as currently recommended in evaluating actual CVD risk.

Treatment with methotrexate reduces the risk for CVD events by ∼20% in RA.20 Indeed, systemic inflammation among patients with RA contributes substantially to increased CVD risk both through its adverse impact on traditional risk factors and direct effects on the vasculature.21 22 Therefore, effective CVD risk management likely comprises not only adequate treatment of conventional risk factors but also tight disease activity control in RA.

With regard to the multiplier application as recommended by the EULAR task force,7 other recent reports revealed that the enhanced CVD risk is unlikely to be restricted to those patients with a disease duration of >10 years, neither to those who experience rheumatoid factor positivity and presumably also not to the small subgroup with extra-articular manifestations.23 The EULAR task force acknowledged that their approach was conservative.3 Taken together, evidence that was mostly reported after the EULAR recommendations for CVD risk management were published raises the possibility that their application could result in a substantial proportion of patients with RA at high risk for CVD remaining unidentified.23 However, are cardioprotective drugs effective in CVD risk management in RA? Indeed, there is a lack of hard evidence regarding CVD prevention in patients with RA and therefore many questions remain unanswered. In this regard, it is both noteworthy and promising that in a post hoc analysis of two prospective trials that assessed the impact of intensive compared with conventional treatment with statins on a CVD outcome, patients with and without inflammatory joint disease were found to experience comparable lipid-lowering effects and reduced CVD...
risk. Also, in a preventive cardio-rheuma clinic, two-thirds of patients referred for CVD risk evaluation required CVD prevention, which further reinforces the need for identifying patients at high risk. RA patients often sustain background marked systemic inflammation, altered lipid parameters and exposure to polypharmacy, all of which can influence optimal lipid-lowering treatment in CVD prevention. Despite the presence of these factors, treatment to lipid targets was successful in as many as ~90% of patients with RA who required interventions with cardiovascular drugs. Experience and studies on clinical CVD prevention are warranted.

Corrales et al. compared the abilities of the mSCORE; carotid ultrasound determined advanced atherosclerosis and CACS in the identification of patients with RA who sustain high or very high CVD risk in the absence of established high or very high risk. Upon using the EULAR task force multiplier, the proportion of patients with high or very high risk increased by only 3.1%, that is, from 11.6% to 14.7%. High CACS (>100) were observed in a mostly similar proportion of all patients, that is, 17.9%, and in none with an mSCORE of <1% (low risk). By contrast, 73.7% of all patients had ultrasonographically confirmed advanced atherosclerosis. Remarkably, this comorbidity was observed not only in 85% of patients at moderate risk (mSCORE>1 and <5%) but also in 33.3% of those with an mSCORE<1%. CACS and ultrasound findings correlated significantly. Nevertheless, even among the 41.2% with no detected coronary artery calcification, 57.5% had carotid plaques. Finally, in keeping with these findings, upon employing the presence of high/very high risk as determined by an mSCORE>5, a CACS of >100 or ultrasonographically determined advanced atherosclerosis as outcome variable, the sensitivities of an mSCORE>5, CACS>100 and ultrasonographically determined advanced atherosclerosis were 19.4%, 23.6% and 97.2%, respectively. The findings on ultrasound in the Corrales study are largely similar to those recently reported by the same group in a larger cohort.

Overall, in the Corrales study, by considering ultrasound findings in addition to the EULAR recommendations, the proportion of patients stratified as being at high or very high CVD risk increased from 22.1% to 77.9% or 3.5-fold. Most importantly, in routine clinical settings these patients would generally not receive adequate preventive CVD risk treatment with the serious consequences this has. Moreover, if up to a third of patients with RA without established high CVD risk and considered to be at low risk are actually at very high risk, it would appear at least reasonable to perform carotid ultrasound not only in those at intermediate risk but also in those with low risk according to the mSCORE. Although the respective proportion was only 13% in the previously reported Corrales study, even the absence of identifiable carotid artery plaque by ultrasound still does not fully exclude the possibility of prevalent significant coronary artery disease.

Could carotid artery plaque associate with lower incident CVD event rates and why would CACS be less sensitive in discerning high risk in RA? Both carotid artery atherosclerosis and CACS predict incident CVD events beyond other risk factors in RA. Vulnerable plaques are more echolucent and typically have a lipid-rich core, macrophages and a low collagen content. By contrast, plaques that are less vulnerable to rupture are more echogenic and contain more collagen, dense fibrous tissue and various amounts of calcification and represent more advanced disease. Patients with RA experience a disease activity-related increased vulnerable plaque burden. It is therefore highly unlikely that reliance on the presence of carotid plaque translates into an overestimate of CVD risk in RA, and it is indeed expected that CACS are less sensitive in this context. The latter is, however, also recognised in non-RA subjects.

Upon considering the potential use of carotid ultrasonography in CVD risk stratification in RA, the following issues are equally relevant. In contrast to determining CACS, ultrasonography is inexpensive, does not require radiation and is considered cost-effective. Could the additional use of biomarkers be preferable to vascular imaging upon evaluating CVD risk in RA? Biomarkers of CVD risk are valuable in examining atherogenic mechanisms in RA and but the identification of those that predict CVD events beyond other risk factors in this disease is in its early stages at present and requires intensive and careful exploration. Also, employing (presumably) a panel of useful biomarkers is likely to enhance the involved costs to a larger extent than performing carotid ultrasonography. Nevertheless, plaque represents advanced atherosclerosis and hence biomarkers that reflect enhanced atherogenesis in RA may well be needed to timely reclassify patients with RA in CVD risk groups and identify those at high risk at a stage prior to plaque occurrence. Finally, whereas plaque associates closely with coronary artery disease, increased carotid intima-media thickness (CIMT) represents mostly high blood pressure-mediated arterial media hypertrophy and relates more strongly to left ventricular hypertrophy. Omission of CIMT results in the Corrales study would not be anticipated to alter the findings as only one of the patients with a CIMT >0.9 mm did not have plaque.

The article by Corrales and colleagues does not allude to the limitations of their investigation that do, however, require further elucidation. Their cross-sectional design precludes drawing inferences on the direction of causality and, accordingly, the role of carotid ultrasonography in CVD risk assessment and management and reduction of cardiovascular event rates needs evaluation in a longitudinal study. Also, since the mean disease duration was 10.8 years it remains to be clarified whether carotid ultrasound is as helpful among patients with early disease versus those with long-standing disease in enhancing CVD risk stratification.

Improved risk stratification alone will not reduce cardiovascular event rates, unless it is accompanied by adequate CVD risk management in RA. Recently reported retrospective data suggest that the latter often does not occur and traditional risk factors are underdiagnosed and undertreated in RA, a situation that could in itself contribute to enhanced CVD risk. This deficiency is amply confirmed in a prospectively designed cross-sectional investigation on CVD risk factor control in 836 patients with RA by Primdahl and colleagues, as also reported in Annals of Rheumatic Diseases. Most striking is that among the 644 patients without established CVD or diabetes, inadequate blood pressure and lipid control were documented in 35.8% and 55.4% of participants, respectively. Even more concerning is that among those with CVD and diabetes, these proportions were as high as 36.2% and 84.2% and 73.6% and 80.9%, respectively. Clearly, effective and innovative measures aimed at improving systematic evaluation and treatment of unfavourable CVD risk factor profiles by healthcare providers in patients with RA are urgently needed. In line with the findings in the Corrales study, upon applying the EULAR task force multiplier, the proportion of patients with high or very high risk increased by only 3.6%, that is, from 12.6% to 16.2%.

Perhaps the most pertinent issue that requires clarification here is: who should
take the responsibility to manage CVD risk in patients with RA? Enrolment in a preventive clinic with cardiologist involvement as currently done in a centre in Oslo, Norway, is likely to constitute one promising option in this regard. However, the establishment of such clinics may not be possible in less well-resourced countries. The Primdahl study investigators fully informed the patient, relevant hospital department and treating general practitioner (GP) about the participant’s risk profile and subsequently referred the patient to their GP. In this regard, in The Netherlands, the GP now assesses and manages CVD risk in patients with RA, a process that is facilitated by embedding of recommendations in the corresponding relevant guideline. Again for patients who form part of less affluent societies, this may not be feasible as it involves extra direct and indirect costs. Should, at least in some settings, the treating rheumatologist manage CVD risk in addition to obtaining optimal disease activity control? At the very least, population studies in some settings, the treating rheumatologist manages CVD risk in patients with rheumatoid arthritis and other forms of inflammatory arthritis. Ann Rheum Dis 2010;69:325–31.

In conclusion, whereas the mortality gap between patients with RA and the general population reportedly continues to widen, considering the findings in the Corrales and Primdahl studies could contribute to the implementation of potentially effective strategies in our attempts at reducing cardiovascular risk in RA.

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