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Association of HbA1c variability with atherosclerosis in diabetes: simple marker, risk factor or statistical bias?

In this issue of the Journal of Diabetes and its Complications, Yang et al. (2015) report that HbA1c variability is significantly associated with the development of subclinical coronary atherosclerosis in type 2 diabetes and that this glycemic disorder is a greater predictor of premature coronary damages than mean HbA1c at early stages of the disease. At first glance, this finding seems to provide an additional contribution to the debate that was opened by Kilpatrick, Rigby, and Atkin (2008). By retrospectively analyzing the data sets of the Diabetes Control and Complications Trial (DCCT) these authors showed that variability in HbA1c adds to the mean HbA1c in predicting the risk of microvascular complications in type 1 diabetes. Such observations were extended to type 2 diabetes by Sugawara et al. (2012), who have established that HbA1c variability predicts the development of microalbuminuria independently of mean HbA1c in type 2 diabetes. However, the investigations of the associations between HbA1c variability and diabetic complications were more often limited to microvascular complications in type 1 diabetes than extended to macrovascular outcomes in type 2 diabetes (Kilpatrick, 2012). Consequently, the report of an increased risk for macro- and micro-vascular events in type 2 diabetes, with worsening HbA1c variability (Hirakawa et al., 2014) is an important finding.

Reverting to the study of Yang et al. (2015), there arises the question as to whether long-term oscillations of HbA1c around a mean HbA1c value can be considered either a simple marker or a causative risk factor of premature atherosclerosis. Unfortunately, the study of Yang et al. (2015), which is observational in design, is unable to answer this question and to eliminate all the other confounding markers or risk factors. The main reason relates to the fact that the multiple logistic regression analysis used in this study is a statistical method, which is mainly designed for establishing associations between a series of “predictors” and a dependent variable (Zar, 1999). In the present study, the dependent variable is coded as either the presence or absence of subclinical coronary atherosclerosis. Apart from the age, which is considered a well-recognized risk factor for harmful vascular outcomes, Yang et al. (2015) have used and selected the following “predictors”: i) those reflecting the overall glucose exposure and glycemic variability ii) those related to lipid disorders, and iii) those concerning specific treatments with either statins or insulin.

As expected, the age and duration of diabetes appear as the best “predictors”, but the results are more questionable when the investigators (Yang et al., 2015) enter mean HbA1c and HbA1c variability as predictive variables using arbitrary models for the statistical analysis. For instance the mean HbA1c never appears as a significant predictor when the HbA1c variability is entered. In contrast, mean HbA1c was statistically associated with the presence of subclinical coronary atherosclerosis when the HbA1c variability was deleted. These somewhat surprising and controversial results are probably due to a bias in the methodology of the multiple logistic regression analysis that requires first eliminating any intercorrelation between the “predictor” variables and secondly avoiding any arbitrary choice for the models selected in this analysis. For instance, it is highly likely that the ambient hyperglycemia and HbA1c variability are positively intercorrelated. In addition, the selection of predictor variables should be done using a stepwise procedure in order to select the model, which is preferable to another. Finally, when we use a regression model, we ideally hope that there is a cause and effect relationship between the dependent variable and the predictive variables (Zar, 1999). Such an idyllic view is rarely confirmed because regression models are more descriptive than predictive. Bringing all these methodological limitations together, we suggest that the issue of the contribution of HbA1c variability to the development of vascular complications, if it exists, cannot be simply solved by a multiple logistic regression analysis and more particularly when such an analysis is affected by statistical bias in methodology.

For that reason, such an issue would warrant to be addressed by implementing an interventional trial with an “optimal” design aimed at either confirming or refuting the aforementioned expected relationship. Ideally, such an interventional trial would involve a randomized controlled study with a long-term follow up period and parallel comparison between two groups of patients with type 2 diabetes, according to the guidelines provided by the Food and Drug Administration, Center for Drug Evaluation and Research (2008). The first one should be assigned to dietary and pharmacological measures aimed at reducing the HbA1c variability to its lower level and, ideally, at achieving a flat time course of the HbA1c profile. The patients assigned to the second group should be maintained on their usual dietary habits and standard antidiabetic treatments. Furthermore, in both groups, the therapeutic strategies should be aimed at achieving similar levels in mean HbA1c, plasma lipids and blood pressure. As it could be difficult to ensure a tight control of all these parameters over several years we should be very cautious about the reliability and pertinence of such long-term studies that have the ambitious goals of testing the impact of HbA1c variability on vascular outcomes, especially when many other risk factors can be involved as potential or real key players in the pathogenesis of diabetic complications (Gaede, Lund-Andersen, Parving, & Pedersen, 2008; Laakso & Lehto, 1997). In addition, the real cause of the improvement in vascular

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outcomes with HbA1c flattening, if demonstrated, might remain uncertain because of the unavoidable lifestyle changes that could act as confounding factors.

Even though we have intuitively the feeling that long-term glycemic variability can exert harmful effects on vascular outcomes, this issue is not likely to be solved in the near future, as the designs of trials able to provide a clear answer are probably too complex and costly. So that the reader does not remain disappointed by the mixed conclusions of the present editorial, we would remind that even in “hard scientific domains” such as mathematics, several decades or centuries are sometimes required to elucidate theorems that could not be definitively demonstrated at the time they were proposed.

References


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