The Obesity Paradox or Vulnerability of the Underweight

To the Editor:

In response to the interesting articles from Zamora1 and Artham,2 we would like to comment on the controversial issue of the obesity paradox.

Advances in pharmacological treatments have been the main reason for the reduction in mortality associated with acute myocardial infarction (AMI) between 1975 and 1995.3 It is therefore surprising that the studies reporting on different aspects of reverse epidemiology, within the context of polypharmacy and haemodynamic instability, as well as acute coronary syndrome (ACS) and acute cardiac failure (ACF), all under-utilise pharmacological variables and are limited to a general analysis of the percentage of drugs used.4,5

Another element to be highlighted is the high percentage of patients excluded from these researches since their weight and/or size is not provided.4 This would seem to suggest that, in particular during the first stages of a cardiological emergency, there are failures in taking the patient’s anthropometric measurements, which may lead to an incorrect pharmacological dose.4

Unfortunately, drugs which have proven to be beneficial in reducing cardiovascular morbimortality can also have very serious adverse effects if the correct dose is not given.3 In this way, subjects with reduced body mass have greater “pharmacological susceptibility,”6 which is perhaps related to older age1,6 and comorbidity, as well as a reduced “therapeutic window.”

Another element to be highlighted is the high percentage of patients excluded from these researches since their weight and/or size is not provided.4 This would seem to suggest that, in particular during the first stages of a cardiological emergency, there are failures in taking the patient’s anthropometric measurements, which may lead to an incorrect pharmacological dose.4

Therapeutic intervals are directly and proportionately related to body mass index (BMI) and are influenced by multiple factors, such as age, sex, and renal function. This relationship is not taken into account in the exclusive analysis of the percentage of drugs used. Fonarow recently recognised the need to include more complex pharmacological variables such as: dose, tolerability and adverse effects.7

It is also to be noted that the insufficiencies indicated have been concluded from studies based on hospital records4,5 and that the situation in practice may be much more critical.

Moreover, markers should not be confused with risk factors.2 The obesity paradox does not fit the causality criteria, since on analysing the strength of the association, severe obesity does not yield better outcomes, in particular when compared to overweight and slightly obese patients.2 While the paradox is not detected or even disappears (“reversal of the reversal epidemiology”)8 in situations where the importance of quantifying the acute pharmacological management is “reduced,” as is the case of sudden death1,2,6 (in particular where this occurs outside the hospital), stable coronary heart disease,9 long-term monitoring (>5 years) of heart failure6,8 or heart failure with ejection fraction >40%,1,2 and heart transplant.8

Finally, we believe that the risk factors can never be separated from the constant of cardiovascular risk, rather they are associated with ACS and ACF in a much more complex maze of causality than we have been able to quantify and in which medical treatment plays a leading role.4

In conclusion, we have put forward a theory that pharmacological variables are the main confounding factors in reverse epidemiology and suggest caution on accepting the validity of the obesity paradox, until more evidence is acquired.

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Response

To the Editor:

We would like to express our thanks to Alberto Morales Salinas for his letter. In our response, we wish to discuss his comments and our manuscript without entering into a debate on the specifics of “reverse epidemiology” described in different pathologic situations. Clearly, in our series incidence of mortality is higher among low weight patients. This is not surprising and there may well be no single factor to justify it. While therapeutic intervals can have a direct, proportionate association with body mass index (BMI) and, in turn, be influenced by multiple factors such as age, gender or kidney function—as Morales Salinas affirms—we are unaware of any study of mortality in heart failure with beta-blockers or ACE inhibitors—the drugs that have most influenced these patients’ survival—that has shown an association between the benefits obtained and dosage adjustment for patient BMI or body surface area. Furthermore, we know that mortality in patients with heart failure has been seen to be related to greater blood concentrations of some drugs. Consequently, although we cannot affirm that some patients may not have exhibited this susceptibility, we do not feel we can consider the greater mortality among low-weight patients in our series may have been favored by a relative excess of the treatment received (“the vulnerability of the low-weight patient”). On the other hand, in no case could we consider this phenomenon a confounding factor to the finding that overweight and obese patients had a better prognosis than normal weight patients did. At no point in our manuscript do we affirm the existence of a causal relation between obesity and better prognosis. We simply state that, in a strict 2-year follow-up, overweight and obese patients presented lower mortality. Nor do we have an explanation for this. Although some series report finding a U shaped mortality curve—ie, greater mortality in patients with outlying weights (low- or over-weight)—and that, therefore, severe obesity did not provide patients with greater protection than being overweight or slightly obese, we did not observe this phenomenon. In fact, as we comment in our study, no patients with morbid obesity (BMI >40) died in the 2-year follow-up although, as we are dealing with a small number of patients, we cannot generalize from our conclusions.

We agree that it is difficult to accept the obesity paradox in patients with heart failure, even though it has been identified in studies with thousands of patients. It would be easier to do so if we could find an explanation as to why this paradox arises that did not need to depend—exclusively—on pharmacologic variables.

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Response

To the Editor:

With great interest, we read Dr Alberto Morales Salinas’s comments in which he questions the paradoxical association between obesity and prognosis in heart failure. Throughout the last decade, numerous cohort studies have been published which detail the so-called “obesity paradox” in the context of both acute and chronic heart failure. Body mass index (BMI) is not the only conventional cardiovascular (CV) risk factor that has a favorable influence in patients with heart failure, given that high concentrations of low density lipoproteins and total cholesterol, as well as high blood pressure, have also been associated with a survival advantage in heart failure.

We agree with Dr Salinas’s ideas and point of view on this controversial topic and recognize the interest of his new hypothesis in as much as most of these studies lack an element of control of the pharmacologic agents used, which could introduce confusion into the final results. We agree that documenting patient height, weight, BMI and kidney function forms part of providing top quality attention and that we need this documentation to avoid dosage errors and unfavorable clinical course.

This new hypothesis may be more applicable to studies conducted in the acute context—when patients are attended while presenting acute decompensated heart failure or acute coronary syndrome—because this is when...
medication dosage tends not to be proportionate to weight, height and BMI. However, even in these acute situations, a vast amount of data points to the existence of an inverse relation between obesity and prognosis. In contrast, the opposite occurs in patients with stable coronary disease (CD), or with other chronic conditions—eg, chronic obstructive pulmonary disease (COPD), rheumatoid arthritis, and terminal kidney disease (TKD)—and the “obesity paradox” phenomenon, or in older patients, in whom this new hypothesis is less likely to play a role in “reverse epidemiology.”

One specific reason that could explain why the “obesity paradox” is observed in such a wide range of diseases—CD, heart failure, arterial hypertension, and dyslipidemia—could be the more energetic treatment administered to obese patients. In one study of patients with CD, the highest BMI values were associated with better administration of CD treatment according to established guidelines, and led to lower rates of in-hospital mortality. Many studies document clear evidence of the fact that low weight patients are not the only ones who present a worse prognosis. Moreover, patients with an ideal body weight—“the golden weight” according to Sarnak—were considered “cachexic.” Logistic regression analysis found the highest percentage of body fat (χ²=9.1; P=.002) was the most powerful, independent predictive factor for illness-free survival. In this population, for every 1% absolute increase in percentage body fat we found a >13% reduction in major clinical episodes. Various possible explanations exist for the inverse association between BMI and mortality: it is crucial to investigate the differences in pharmacologic agent dosage, secondary effects and tolerability in relation to BMI. This could provide a partial clue to the explanation and, therefore, we think there is a clear need for new clinical studies to clarify fully the mechanism underlying these paradoxical relations, in the hope they lead to new, definitive treatments.

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