In the previous issue of Critical Care, Wang and colleagues [1] present interesting data from a large cohort of unselected medical intensive care unit (ICU) patients which examined the prognostic utility of two well-established biomarkers: N-terminal pro-brain natriuretic peptide (NT-proBNP) and C-reactive protein (CRP). In fact, the authors’ observations nicely complement the picture that is emerging from several recent studies [1-14]. Like most of the previous studies, their findings leave the majority of ICU physicians in doubt about whether biomarkers are utile or futile.

The ICU is a rather hostile setting for biomarkers. Biomarkers complement other clinical information by proving quantitative data regarding a pathophysiological mechanism that can be used for the early diagnosis of a specific disease, to monitor and guide treatment, and to predict the risk of death or other adverse events. The stronger the link between the information provided by the biomarker and the immediate clinical course of action that we physicians take in response, the higher the clinical utility of the biomarker [13,14].

In most patients finally being admitted to an ICU, the diagnosis is made prior to ICU admission, most commonly in the emergency department (ED). Of course, we still face diagnostic uncertainty in many patients who develop new symptoms or signs during their stay in the ICU (for example, respiratory deterioration or fever). To appropriately examine the diagnostic accuracy of a biomarker in these settings, we need to define a gold standard diagnosis against which the blinded biomarker results are then compared. Unfortunately, owing to, for example, the low specificity of chest x-ray findings, the adjudication of a final diagnosis often is challenging for many common ICU disorders, such as ventilator-associated pneumonia and hypoxic respiratory failure [7,10,12]. In addition, the extent to which experience and diagnostic cutoff levels can be transferred from studies performed in the ED to the critically ill patients in the ICU is questionable [7,10-14]. Major differences in patient characteristics, disease severity, comorbidity, resources available for the individual patient, and therapies applied between the ICU and the ED require that the potential clinical use of biomarkers in the ICU be defined by specific ICU studies.

What about the utility of biomarkers in monitoring treatment? Biomarkers are used routinely to monitor the efficacy and safety of treatment. For example, urine output and serum creatinine are used to quantify renal function; tidal volumes, oxygen saturation, and arterial partial pressure of oxygen [PaO₂] are used to tailor ventilator settings; and body temperature, CRP, and procalcitonin are used to assess the response to antibiotics. Although the use of biomarkers in many of these indications is mainly empirical and only partly supported by large prospective studies, it is perceived by most clinicians as utile as the links between the biomarker information and therapeutic consequences are strong [11].

The case is more challenging for prognostic biomarkers. The link to a specific consequence is weakest for...
prognostic biomarkers applied in patients with a wide variety of diseases, such as in unselected ICU patients. The added value of most, if not all, previously examined biomarkers on top of current ICU mortality scores seems to be too low to justify clinical use [1-6]. The prognostic accuracy for ICU or in-hospital death of most biomarkers is modest and inferior to that provided by, for example, the APACHE (Acute Physiology and Chronic Health Evaluation) score [1-7]. This observation seems to be well explained by the wide range of disorders leading to ICU admission and the fact that different organ systems may be the most severely damaged and therefore critical for survival. Moreover, it is important to highlight that there is no perceived unmet clinical need to appropriately risk-stratify most patients in the ICU. Simple clinical variables, many of which are captured in the ICU scores, provide immediate and reasonable risk prediction. As cardiovascular function is the key variable in many critically ill patients, BNP and NT-proBNP – as quantitative markers of hemodynamic cardiac stress and heart failure summarizing the extent of systolic and diastolic left ventricular dysfunction, valvular dysfunction, and right ventricular dysfunction – have been shown to be predictors of death in several previous studies. We are still searching how to best apply this information in the clinical care of critically ill patients.

However, prognostic biomarker studies, particularly with BNP and NT-proBNP, have already contributed to a better understanding of many disorders in the ICU. In fact, the observation that hemodynamic cardiac stress is present in multiple conditions provided important novel insights into pathophysiology and highlighted a dominant role of the cardiovascular system of many common disorders in the ICU, including septic shock and weaning failure [8,9,12]. Ultimately, these insights will contribute to improvements in our management of ICU patients.

Abbreviations
BNP, brain natriuretic peptide; CRP, C-reactive protein; ED, emergency department; ICU, intensive care unit; NT-proBNP, N-terminal pro-brain natriuretic peptide.

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