jaundice, in which oxidative hemolysis may play an important role, is more common among newborns with the *A/*C genotype than among those with the *A/*A genotype.

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CSF and G-CSF on these parameters; and (3) the relationship of lactoferrin (Lf) plasma levels with the initial G-CSF peak and the subsequent neutrophil rescue. Modifications of plasma cytokine levels may be due to changes in production, release, consumption, or clearance. This clinical approach allows to establish temporal correlations between hematopoietic growth factor (HGF) administration and fluctuations of endogenous serum cytokines. However, the underlying cause/effect mechanisms remain largely hypothetical. Furthermore, because exogenous GM-CSF or G-CSF was combined with Epo, the specific role of each cytokine cannot be determined. Despite these limitations intrinsic to the clinical setting, the results provide novel information on the interaction between exogenous HGFs and the SCT-related cytokine release network.

We have also monitored the plasma concentrations of HGFs (ie, IL-3, GM-CSF, G-CSF, Epo, IL-8, IL-6, and IL-5) and granule proteins (Lf and myeloperoxidase [Mpo]) in 21 ovarian cancer patients undergoing autologous PB SCT (Table 1). G-CSF levels were markedly higher after exogenous G-CSF/Epo, as expected; furthermore, they were significantly lower following GM-CSF/Epo than after SCT alone; GM-CSF levels, moderately elevated after exogenous GM-CSF/Epo, were similar in the G-CSF/Epo and control groups. Finally, patients treated with either G-CSF/Epo or GM-CSF/Epo showed higher Epo plasma concentrations than the group receiving no exogenous HGFs (Fig 1 and data not shown).

When compared with SCT alone, (1) G-CSF/Epo treatment moderately increased the IL-3 level peaking at day +10, and did not significantly modify M-CSF level (data not shown), whereas it almost completely suppressed IL-8 and moderately decreased IL-6 and IL-5 levels (Fig 1); (2) GM-CSF/Epo treatment increased M-CSF, associated with a moderate increase of IL-3 levels peaking at day +10, moderately decreased both IL-5 and IL-8 concentration and did not modify IL-6 (Fig 1 and data not shown).

These observations indicate that exogenous G-CSF treatment induces a marked decrease of IL-8 levels and moderately decreased IL-6 concentrations, whereas exogenous GM-CSF markedly reduces G-CSF and IL-8 concentrations and increases M-CSF levels. The increase of IL-3 and the decrease of IL-5 levels were similarly observed in patients treated with G-CSF/Epo or GM-CSF/Epo. Altogether, these results indicate that exogenous HGFs modulate endogenous cytokine levels; consideration of these aspects may contribute to optimize HGF treatment protocols in the clinical SCT setting.

We also evaluated the relationship between these phenomena and Lf release: (1) in all three groups the Lf concentration is strictly and directly correlated with the neutrophil response with respect to time-response patterns and plasma levels. (2) In patients subjected to SCT alone, the G-CSF decrease after the G-CSF peak coincides with initiation of the Lf response; the inverse correlation between these two parameters is highly significant (data not shown). Altogether, these temporal correlations after SCT provide circumstantial evi-
dence that, after the G-CSF peak, the generated granulocytes release Lf, which could negatively feedback on the release of G-CSF, thus in line with the experimental model proposed by Broxmeyer.15

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