Biomarkers in Transplantation Medicine: Prediction of Pharmacodynamic Drug Effects

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Imunosuppressive drugs · Whole blood · Pharmacodynamics · Flow cytometry · T cells · Dendritic cells · Biomarkers · Transplantation

Summary
Conventional therapeutic drug monitoring based on measuring of blood concentrations (pharmacokinetic) is important in the clinical management of immunosuppressive therapy in transplantation medicine. Since rejection or infection occurs at irregular drug concentrations, immunosuppressive drug therapy is often empiric and prophylactic in nature. In addition, blood immunosuppressant levels are only indirect predictors of the pharmacologic effects on immune cells (pharmacodynamic) because, due to the genetic heterogeneity, the immune systems of the transplant recipients are not equally sensitive to drug effects. Therefore, therapeutic drug monitoring requires the application of reliable and effective methods to study the pharmacodynamic variability by direct measurements of drugs effects on immune cell functions. Against this background we developed assays which are based on whole blood, flow cytometry and biomarkers of diverse functions of T cells and of dendritic cell subsets. These biomarker assays allow us to differentiate between synergistic and antagonistic pharmacodynamic effects of an immunosuppressive drug combination therapy in vitro and to predict the pharmacodynamic drug effects in heart-transplanted recipients. Such a pharmacodynamic drug monitoring based on biomarkers may offer the opportunity to complete conventional therapeutic drug monitoring and, therefore, to tailor immunosuppressive therapy more individually.
Introduction

Despite the introduction of new sampling strategies for measuring drug concentrations and despite the use of new potent agents, therapeutic drug monitoring after organ transplantation is still challenging in the clinical routine. The small therapeutic window of all currently available drugs requires a monitoring which is done by routinely measurements of blood or plasma drug concentrations (pharmacokinetic) to avoid toxic drug levels and to monitor recipients’ compliance. However, the limitations of the pharmacokinetics to measure pharmacodynamic drug effects has prevented immunosuppressive therapy from being administered at high levels ‘pre-emptively’ for those recipients who require more intense immunosuppression, leading to rejection and increased risk of late graft failure [1]. In addition, combinations of immunosuppressive drugs are associated with synergistic and antagonistic effects. Moreover, inter-subject variability may occur with respect to the sensitivity to suppression of immune function. Thus, in order to balance immunosuppression, one current goal of transplantation medicine is to explore peripheral biomarkers of relevant immune events to monitor the immunosuppressive status of recipients for prediction of the pharmacodynamic drug effects [2].

In predictive medicine [3, 4] the potential of biomarkers for cardiovascular diseases and clinical outcome after cardiac surgery with cardiopulmonary bypass have been investigated [5–7]. In earlier studies [8–10] we evaluated the potential of biomarkers of T cell functions in an experimental animal model. We determined the relationship between the pharmacodynamic effects and both the pharmacokinetics and the drug doses. Furthermore, we observed that the changes of T cell function over time in immunosuppressed allograft recipients correlated highly with the severity of histopathologic events within the allograft tissue [8]. In addition, we detected new mechanisms of drug action and could distinguish between synergistic, additive or antagonistic drug interactions after combination therapy [9, 10].

This article highlights our results of using biomarkers of T cell and dendritic cell functions in human in vitro studies and of clinical studies in heart-transplanted (HTx) recipients in the context of the current literature.

Methods

Our pharmacodynamic assays rely on the following methods: i) whole blood as the matrix of choice to reflect the effects of immunosuppressive drugs in vivo; ii) flow cytometry analysis; iii) biomarkers of immune cell functions.

We used whole blood instead of the generally used purified cells or isolated cell lines as the matrix for our assays for various reasons [11]: i) the use of whole blood requires 10- to 100-fold lower sample volumes and shorter preparation times compared to techniques that rely on purified cells or cell lines; ii) nonuniform cell loss during separation alters cell populations and disturbs cell-cell interactions; iii) whole blood includes plasma components and red cells to which drugs bind and into which they distribute; iv) most important, a dissociation of drug from its cellular receptor may occur during the process of cell purification; v) a more frequent blood sampling is possible because of the required low amount of blood.

The amount of whole blood in our assays varied from 2.5 ml for in vitro studies (various drug concentrations) to 20–200 µl, depending on the investigated immune function, for the ex vivo studies for a single time point. We used nonspecific stimuli to activate immune cells rather than specific stimuli, e.g. allogenic spleen cells, because the latter have the disadvantages of being not available for repeat testing and of stimulating only specific immune cell clones [12]. However, triggering of selected immune cell activation pathways is important for a better understanding of the mechanisms of action of immunosuppressants. This allows to devise immune function assays which enable to distinguish drug effects [13]. Thus, we used the following mitogens to assess immune cell functions: i) concanavalin A: T cell proliferation and surface antigen expression; ii) phorbol 12-myristate 13-acetate plus ionomycin: intracellular T cell cytokine production; iii) actinomycin-D: T cell apoptosis induction; iv) lipopolysaccharide (LPS) (Escherichia coli): intracellular dendritic cell cytokine production.

First, we studied the changes of immune cell functions over time in whole blood stimulated with different mitogen concentrations to define optimal assay conditions. Second, we used optimized assays in vitro to evaluate the drug potencies and efficacies and to differentiate between synergistic and antagonistic effects by adding different drug concentrations alone or in combination to whole blood. Third, in clinical studies we tested the immune function assays to assess the pharmacodynamic drug effects of treated transplant recipients by ex vivo stimulation of whole blood.

Flow Cytometry

It has been proposed that the flow cytometric investigation of whole blood cell cultures is particularly suitable to observe the pharmacodynamic drug effects with high sensitivity, versatility, and speed [14, 15]. It can also be used for in vitro testing [16, 17]. In comparison to techniques relying on radionucleotide incorporation to measure inhibition of cell proliferation as an index of drug effect, flow cytometry has been shown to be more specific in measuring drug potencies and efficacies in vivo [18]. Moreover, flow cytometry allows the simultaneous detection of various parameters of immune functions in a small sample volume with a high throughput of cell analysis.

Immune Cell Functions

T cell proliferation was measured by simultaneous analysis of proliferating nuclear antigen (PCNA) and DNA content,
Table 1. Characteristics of cellular differentiations (CD) of T cell surface antigens

<table>
<thead>
<tr>
<th>CD antigen</th>
<th>Function</th>
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</thead>
<tbody>
<tr>
<td>CD11a (LFA-1 α-chain)</td>
<td>adhesion (binding (ICAM-1)</td>
</tr>
<tr>
<td>CD25 (IL-2 γ-chain)</td>
<td>cell proliferation</td>
</tr>
<tr>
<td>CD71 (transferrin receptor)</td>
<td>induction of apoptosis</td>
</tr>
<tr>
<td>CD95 (Fas antigen, APO-1)</td>
<td>adhesion, co-stimulation</td>
</tr>
<tr>
<td>CD134 (OX40)</td>
<td>induction B-cell proliferation and activation after T-cell contact</td>
</tr>
<tr>
<td>CD154 (ligand of CD40)</td>
<td></td>
</tr>
</tbody>
</table>

LFA = Leukocyte function antigen; ICAM = intracellular adhesion molecule.

whereas T cell activation was detected by the expression of surface antigens playing a potential role in the immune response, e.g. co-stimulation (CD25, CD71), adhesion (CD134, CD154) or apoptosis (CD95) (table 1). Furthermore, T cell activation was detected by the accumulation of intracellular cytokines such as IFN-γ, TNF-α, IL-2 and IL-4 [7, 19]. Additionally, T helper 1 (TH1) and TH2 cytokines (IL-2, IL-4, IL-6, IL-10, TNF-α and INF-γ) were detected in the serum using multiplex bead array analysis [20].

Apoptotic T cells were quantified using the fluorescent annexin V, and the TUNEL assay was used to detect DNA breaks [21].

The myeloid and plasmocytoid dendritic cell subsets in whole blood were determined without stimulation by specific gating strategies to distinguish these two lineages from leukocytes [22]. However, the dendritic cells’ production of the intracellular cytokines IL-1β, TNF-α, IL-6, IL-8 and IL-12 was measured after LPS stimulation of whole blood [23].

Results

Current practice of immunosuppressive maintenance therapy after organ transplantation consist of combining one basis drug with one adjunctive drug, which have different mechanisms of action during the cell cycle and each bind to a specific target enzyme [24]:
- G0 phase: cyclosporine (CsA) and tacrolimus (TRL), both calcineurin inhibitors;
- G1 phase: sirolimus (SRL) and its derivate everolimus (ERL), both inhibitors of mammalian target of rapamycin;
- S phase: mycophenolate mofetil (MMF), the pro-drug of the active metabolite mycophenolic acid (MPA), inhibitor of ionosine monophosphate dehydrogenase.

Therefore, different drug combinations are possible: CsA and TRL are basis drugs; SRL and ERL are basis drugs as well as adjunctive drugs, and MMF is an adjunctive drug.

In vitro Studies

In general, assessment of drug effects depends on various factors such as cell type, cell activation status and proliferation and/or apoptosis status and on the assay itself. Nevertheless, our apoptosis assay provided insights into mechanisms of drug monotherapy. Thus, we detected a stronger effect on apoptosis in T cells for MPA compared to SRL or TRL, whereas CsA has no effect [21], confirming a previous study [25].

Evaluating the effects of different drug combinations we showed that overall MPA plus TRL had a greater impact on inhibition of T cell proliferation, and to a lesser extent on T cell surface antigens, than all other drug combinations [13].

Our data confirmed that the contribution of MPA is through its substantial antiproliferative effect, but, our data also suggested that MPA mediates its suppression by inhibition of the expression of several T cell surface antigens as earlier proposed by our study group based on an experimental model [8].

Thus, MPA increased the TRL effect on surface expression inhibition of T cell activation antigens in a independent synergistic way. Interestingly, the effect of MPA, regardless of the concentrations used, on T cell surface antigens (CD71, CD95, CD134, CD154) in combination with CsA was antagonistic despite an independent synergistic effect of both drugs at the lowest concentrations on CD25.

In contrast to MPA, the effect of SRL in combination with either CsA or TRL on both PCNA and surface antigen expression was initially synergistic but with increasing concentrations antagonistic effects were observed which occurred at the same time in experiments with partial agonists [13]. This is in line with results of studies using human peripheral blood mononuclear cells or whole blood [26, 27].

Even though the differences in the mode of action of these drugs during the cell cycle might satisfactorily explain our results, limitations of drug actions such as competitive binding to the specific immunophilin or a varying active amount of the immunophilin [28, 29] might contribute to our findings, too.

Clinical Studies

Earlier biomarker assays measured drug target enzyme activity in renal transplant recipients to predict efficacy of CsA [30] and SRL [31] or to evaluate the predictive value of pre-transplant values of MPA enzyme inhibition for the post-transplant outcome [32]. The enzyme assays, in general, do only measure enzyme activity, and not the numerous effects on immune cell function caused by enzyme inhibition. Furthermore, enzyme assays do not measure the net effect on the immune system caused by multiple immunosuppressive drugs with different mechanisms of action. In addition, up to 20 ml of whole blood are required to perform the enzyme assays which restricts its use for repeat measurements or in pediatric recipients.

A commercially available assay measuring ATP levels has been used to investigate the correlation between ATP levels and clinical outcome in renal transplant recipients [33]. Despite the potential of ATP levels to be an early biomarker of
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efficacy, it reflects the overall status of cell metabolism rather than to determine specific insights to drug effects. T cell cytokine expression in the serum, intracellularly or at messenger RNA levels, have been evaluated as biomarkers for pharmacodynamic drug measurements in various human transplant studies using ELISA, flow cytometry or PCR as analytic tools [30, 34–40]. The results of these studies differed regarding the relation between the drug pharmacokinetics and drug pharmacodynamics as well as regarding the specification of one to the other cytokine to measure the pharmacodynamic drug effect. This is in line with results of our study about the conversion of a initial immunosuppression of CsA by TRL or SRL [41]. We found that inhibition of cytokine expression correlated with the whole blood TRL concentrations, whereas in the SRL group the increased blood SRL concentration reflected an increase of cytokine inhibition only on day 4 after conversion. The least effect on expression in our study, regardless of the basis immunosuppressive drug used, was detected for IL-4 when compared to inhibitory effects on IL-2, TNF-α and IFN-γ expression [41].

In addition, using the multiplex analysis for simultaneous detection of several TH1/TH2 cytokines, we observed a significant decrease of cytokine expression (TNF-α, IFN-γ, IL-2 and IL-6) 2 h after drug dosing. These drug effects were reflected by significantly increased CsA and MPA concentrations 2 h after treatment whereas in unstimulated whole blood no drug effects on cytokine production could be observed, underlining the above mentioned advantages of mitogen stimulation of whole blood for quantitating suppression of immune cell function [8, 36].

Because of the circadian cycle of cytokine expression, it was suggested to conduct drug application and blood sampling in clinical pharmacodynamic studies at specific time points to avoid that these circadian fluctuations falsify the monitoring of cytokine expression [12]. It was proposed that the pharmacodynamic parameter expression over time might be more appropriate to measure the drug’s pharmacodynamics than the measurement at one single time [8, 36]. Due to the difficulties to obtain blood at several time points, especially in transplant recipients, blood should be analyzed at least at two time points (e.g. before and 2 h after therapy) to provide information about drug efficacy and about the relationship between pharmacodynamics and pharmacokinetics [42].

In a study in stable HTx recipients treated with a combination therapy of CsA plus MMF, we obtained a high correlation between CsA effects and both blood CsA concentrations and CsA dose. A significant increase in blood CsA concentration 2 h after dosing led to a significant decrease in PCNA as well as T cell surface antigen expression [41]. Furthermore, it could be shown that CsA blood concentrations were more strongly correlated to T cell proliferation and surface antigen expression than to T cell cytokine release. This might implicate to prefer PCNA and surface antigens to cytokines as biomarkers in the combination therapy of CsA and MMF. Additionally, the early antiproliferative effect of CsA might be due to additive effects in combination with the antiproliferative drug MMF as observed in our earlier studies [10, 13]. Interestingly, another of our HTx studies [43] showed contradictory results in this study the increased drug concentrations 2 h after application of either CsA plus MMF or TRL plus MMF did not further attenuate the apoptotic T cell rate, suggesting a steady state drug effect on T cell apoptosis.

The potential of T cell surface antigens as biomarkers to predict the grade of rejection or to predict the pharmacodynamic drug effect was confirmed by other human transplant studies [35, 44, 45]. In our conversion study the increased blood TRL concentrations over time do not further inhibit expression of CD25 and CD95, but significantly inhibit PCNA and surface antigen CD134 expression from the 3rd day after conversion. Another study in renal transplant recipients showed that, after conversion from CsA to TRL, TRL has a significantly higher impact on expression of T cell surface antigens than CsA therapy [46].

After conversion from CsA to SRL the blood SRL concentration increased on day 1 and remained stable at days 3 and 4. But expression of PCNA is not altered by SRL, whereas the expression of CD25 is increased [41]. Our data suggest that SRL dose adjustment relying on blood SRL concentrations do not predict SRL effects on immune cells. This is in accordance with results of our experimental animal study where higher SRL doses led to an increase of blood SRL concentrations without any further inhibition of T cell proliferation or surface antigens [9].

Over the recent years the importance of dendritic cells for the graft outcome has been realized, determining dendritic cells as ideal therapeutic targets for pharmacological modulation to improve allograft survival [47]. Therefore, we directly enumerated two subsets of dendritic cells (myeloid and plasmocytoid dendritic cells) in the peripheral blood of chronically immunosuppressed HTx recipients.

One characteristic feature of all mature but tolerogenic dendritic cells is their low or absent production of proinflammatory cytokines such as IL-12. Long-term liver transplant recipients with or without low-dose immunosuppressive monodrug therapy had a higher plasmocytoid/myeloid dendritic cell ratio than those with standard immunosuppressive therapy [48]. In our study HTx recipients had a lower subset ratio of plasmocytoid to myeloid dendritic cells than healthy human controls, but after conversion from either CsA/TRL or SRL to ERL the plasmocytoid/myeloid dendritic cell ratio increased [49]. Despite their key role to induce tolerance, plasmocytoid dendritic cells also initiate and maintain the antiviral immune response. Thus, cytomegalovirus infection impairs the immunophenotype and the status of maturation of dendritic cells of HTx recipients [50, 51]. In contrast myeloid dendritic cells are bacteria-activated and mainly produce IL-12 to stimulate...
CD4+ T cells [52]. After LPS stimulation of whole blood of recipients treated with ERL, we found a decrease of IL-12 expression compared that detected in recipients with CsA/TRL or SRL therapy [49].

Conclusion

For therapeutic drug monitoring biomarkers can be used to predict pharmacodynamic effects of immunosuppressive drugs. However, until now it is not known to what extent biomarkers of immune cell function may be altered by infection or by rejection. Furthermore, it is not clear whether or not baseline values for individual recipients are required nor is an optimal pharmacodynamic target range known [2]. In spite of the above mentioned advantages of whole blood flow cytometry for the assessment of biomarkers of immune cell functions [16], it is foreseeable that further progress in the prospering field of multiplex flow and slide-based cytometry [53–55] will permit to yield more information on more biomarkers of many more cellular functions in different immune cell lineages from smallest amounts of whole blood [56] and to predict pharmacodynamic drug effects more precisely than today [57–59]. Additional future perspectives are based on diagnostic pharmacogenetic assays to identify inherited genetic factors having an impact on the drug response, on the relative susceptibility to drug-drug interactions, or on the predisposition for side effects [60].

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