Relationship of Microparticles to Progenitor Cells as a Measure of Vascular Health in a Diabetic Population

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Objective: Quantitative measures are needed to identify diabetic patients at higher risk for CV events. Cell-derived microparticles (MPs) are submicron membrane vesicles released from activated cells that are indicative of cell damage. Progenitor cells (PCs) including proangiogenic cells (PACs), often termed endothelial progenitor cells (EPCs), are mediators of reparative capacity. We examined whether the relationship of MPs to PCs/PACs could be used as an improved and clinically feasible index of vascular pathology.

Methods and Results: Plasma samples were collected from patients with early-stage (ES, Diagnosis < 1 year) and long-term (LT, Diagnosis > 5 years) Type 2 diabetes and compared with age related healthy subjects (H). PC and MP subtypes were measured by a combination of flow cytometry and ELISA-based methods. The ratio of procoagulant MPs/CD34+ PCs proved a valuable index to distinguish between subject groups (P = 0.01). This index of compromised vascular function was highest in the LT group despite intensive statin therapy and was more informative than a range of soluble protein biomarkers.

Conclusions: This is the first report of a relationship between MPs and PCs in Type 2 diabetes. This ratio may provide a quantitative and clinically feasible measurement of vascular dysfunction and cardiovascular risk in patients with diabetes. © 2010 Clinical Cytometry Society

Key terms: endothelial; progenitor; vascular; diabetes; microparticles
In this regard, cell-derived MPs and progenitor cells (PCs), defined as CD34 positive cells, and proangiogenic cells (PACs), also CD34 positive along with having endothelial surface markers, have emerged as surrogate markers for cellular damage and repair, respectively. Both have been associated independently with a range of vascular diseases including diabetes (4–7).

MPs are membrane encapsulated vesicles of ~1 μm or less in diameter, derived from surface membranes under conditions of cellular activation or apoptosis (8). These submicron vesicles are released into circulation, carrying with them an array of surface markers, used to identify their cellular source. Exposed membrane phosphatidyl-serine (PS) and tissue factor, along with a plethora of other surface molecules and cytoplasmic components, including nuclear material (9), enable MPs to impact on a variety of biological functions, including coagulation, thrombosis, and angiogenesis (10). Direct receptor-ligand interactions (11) and fusion and delivery of intracellular contents (9,12) are mechanisms employed by MPs to target and impact host cells. Increases in endothelial MPs (EMPs) (13), platelet MPs (PMPs) (14), and monocyte MPs (MMPs) (15) have all been implicated in vascular pathology, including myocardial infarction. Circulating EMPs, generated from the surface of the endothelium have received considerable attention as a surrogate marker for endothelial dysfunction (4,16,17). Because of disease-specific changes and associated biological activity, these vesicles are considered active paracrine agents that aggravate further vascular dysfunction (18) and promote inflammation (19). To date, studies assessing MPs have relied on flow cytometry methods performed in specialized laboratories. The utility of a plate based method has not been tested in comparison to flow cytometry assay.

PACs (formerly given the designation of endothelial progenitor cells) play a significant role in vascular homeostasis, and in the development and resolution of cardiovascular pathologies. For example, the number and function of PACs are altered immediately after a myocardial ischemic event, and in individuals at high risk of pathologic cardiovascular events (7). In addition, MPs are lowered and PACs are increased after treatment with HMG CoA reductase inhibitors (statins) (20,21). A study responding that a prior family history of cardiovascular disease, 12 of the 18 (66%) subjects in the healthy group, 10 of the 11 (90%) in the early stage and every participant (100%) in the long-term group responded that a prior family history of cardiovascular disease existed. The average length of diabetes in the LT patients was 20 years. All study participants were non-smokers. Written informed consent was obtained from all study participants and study protocols were approved by the Institutional Review Board (IRB). Each subject donated ~50 mL of venous blood at ~8 am in the morning and all subjects had fasted the night beforehand. Blood was collected in heparinized (Baxter) syringes for cell and soluble protein analysis (30 mL), Sodium-Citrate for microparticles (10 mL) and EDTA for lipid analysis (10 mL). Demographic data, medical/medication history, physical examination, and vital signs were recorded for each subject.

**Flow Cytometry for PCs/PACs**

Less than 1 h post sample collection, white blood cells were isolated from 30 mL of blood using ammonium chloride lysis as previously described (5). Platelet counts were not determined. Cell staining, gating strategy, flow cytometric methods, and analysis were followed as described (5). Approximately 5E6 cells were stained with a 6-color antibody panel: FITC-anti-CD31 (PECAM) (Pharminingen), PE-anti-CD133 (Miltenyi Biotech.), PerCP-Cy5.5-anti-CD5, CD19,-CD33 (Becton Dickinson), APC-H7 anti-CD45 (Becton Dickinson), PE-Cy7-anti-CD34 (Becton Dickinson), and APC-anti-VEGF-R2 (R&D Systems). Viability was assessed by propidium iodide exclusion. Using a Becton-Dickinson LSRII cytometer, 2E6 live events were processed for each sample and the six fluorescent markers along with light scatter allowed only viable, low to medium side scatter. Singlets that were CD3,19,33-negative were analyzed for PCs and PACs. Singlets were gated as the prominent cluster of cells identified from a plot of side scatter width versus forward scatter width to ensure that cell aggregates

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**RESEARCH DESIGN AND METHODS**

**Patient Recruitment and Study Design**

Patients diagnosed with Type 2 diabetes within the preceding 12 months were termed “Early Stage” (ES) and those diagnosed more than five years ago were termed “Long Term” (LT). Age related “Healthy” (H) subjects were recruited into the study on the basis of having no prior or current history of diabetic- or cardiovascular-related conditions and were not taking any type of CV-related medication including statins or medication for hyperlipidemia, hypertension or diabetes. Four (36%) of the ES group and 14 (67%) of the LT group were receiving statin medication, along with a range of other medications to treat diabetes, hypertension, and other conditions. Regarding family history of cardiovascular disease, 12 of the 18 (66%) subjects in the healthy group, 10 of the 11 (90%) in the early stage and every participant (100%) in the long-term group responded that a prior family history of cardiovascular disease existed. The average length of diabetes in the LT patients was 20 years. All study participants were non-smokers. Written informed consent was obtained from all study participants and study protocols were approved by the Institutional Review Board (IRB). Each subject donated ~50 mL of venous blood at ~8 am in the morning and all subjects had fasted the night beforehand. Blood was collected in heparinized (Baxter) syringes for cell and soluble protein analysis (30 mL), Sodium-Citrate for microparticles (10 mL) and EDTA for lipid analysis (10 mL). Demographic data, medical/medication history, physical examination, and vital signs were recorded for each subject.
were excluded from analysis. Fluorescence minus one (FMO) samples were used as negative controls. Cell populations (PCs and PACs) were quantified as a percentage of mononuclear cells calculated as number per ml blood. Analysis focused on subset definitions for PCs and PACs (Table 1). Data analysis was performed using FlowJo analytical software (Treestar, Ashland, OR).

### MP Isolation

Platelet-poor plasma (PPP) was obtained from citrated blood within an hour after blood collection in order to isolate MPs. Whole blood was centrifuged at 1,500 g for 15 min, supernatant collected, and PPP obtained by centrifugation at 13,500 g for 5 min at room temperature. For each subject, PPP was aliquoted into separate tubes and stored at −80°C until subsequent use. All samples used were subjected to only one freeze-thaw cycle.

### Flow Cytometry for MPs

For characterization and quantification of MPs, PPP was incubated with a mixture of Annexin-V (FITC), PE-CY5-CD41a, APC-CD14 (BD Biosciences) and PE-CD144 (R&D System) in 1X BD annexin-V binding buffer (10 mM Hepes, pH 7.4, 140 mM NaCl, and 2.5 mM CaCl₂) (BD Biosciences) for 30 min at RT in darkness, then 1X BD annexin-V binding buffer was added to make total volume of 1 mL/tube. The negative control was prepared as PPP stained with Annexin-V (FITC) and same amount of matched isotype control antibodies in calcium-free binding buffer. Using a BD Biosciences FACSCanto cytometer, a P1 region (<1 μm) on FSC-H and SSC-H scatter (log scale) was defined by calibrator beads (Fig. 1A). The number of MPs per μL was determined using the P1 region and also 6-μm microsphere beads (Bacteria Counting Kit, Invitrogen) to determine volume of sample (μL) analyzed (Fig. 1B). The number of MPs stained with each specific Ab and Annexin-V was analyzed and determined using FACSDiva software (BD Biosciences) and expressed as MPs/μL. Characterization of

![Fig. 1. Gating strategy for MP analysis. MPs were identified by first gating on the P1 region on the FSC/SSC plot defined by calibrator beads of less than 1 μm (A and B). The origin of the microparticles was determined by coexpression of Annexin-V and CD144 for endothelial derived MPs (C), Annexin-V and CD41 for platelet-derived MPs (D), Annexin-V and CD14 for monocyte-derived MPs (E).](image-url)

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**Table 1**

<table>
<thead>
<tr>
<th>Cell/Particle Type</th>
<th>Marker definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Progenitor cell (PC)</td>
<td>CD133⁺, CD34⁺, CD133⁻/34⁻</td>
</tr>
<tr>
<td>Proangiogenic cell (PAC)</td>
<td>CD133⁺ VEGF-R2⁻, CD34⁺ VEGF-R2⁻, CD133⁻CD34⁺ VEGF-R2⁺</td>
</tr>
<tr>
<td>Endothelial microparticle (EMP)</td>
<td>CD144⁺</td>
</tr>
<tr>
<td>Platelet microparticle (PMP)</td>
<td>CD41⁺</td>
</tr>
<tr>
<td>Monocyte microparticles (MMP)</td>
<td>CD14⁺</td>
</tr>
<tr>
<td>Phosphatidylserine⁺ (PS⁺) MP</td>
<td>AnnV⁺</td>
</tr>
</tbody>
</table>
cellular origin of MPs by positive antibody staining is listed in Table 1.

**MP Plate-Based Capture Assay**

Concentrations of PS$^+$ MPs were determined using the Zymuphen MP Activity Kit (Aniara-CAT#A521096). In this assay, MPs were captured from PPP onto insolubilized annexin-V and their PS content was measured by a functional prothrombinase assay. This offers an indirect measurement of total MP procoagulant activity via measurement of nM of phosphatidylserine on the outer membrane surface. MPs measured by this method are expressed in nM of phosphatidylserine equivalent.

**Soluble Proteins**

Soluble proteins were determined from citrated plasma by electro-chemiluminescent detection using commercially tested kits, as per the manufacturer’s instructions. Meso-Scale Discovery multiplex kits, including the Vascular Injury II assay kit (CAT# K11136G-1) was used to measure SAA, CRP, VCAM1, and ICAM1. IL6, IL8, TNFα, and IL1β were measured using the Human Pro-Inflammatory Base Kit (K15025A-5). The Human Hypoxia Assay (CAT# K15122C-1) measured VEGF, IGFBF-1 and EPO. Plasminogen activator inhibitor (PAI-1) was detected with the Imubind kit from AmericanDiagnostica. Stromal cell-derived factor 1 (SDF-1) was measured by R&D systems (CAT# DY350). IL1β, SAA, IL6, and IL8 were below detection with these assays.

**HbA1c**

HbA1c was performed using the Primus boronate affinity HPLC method (Primus Corporation, Kansas City, MO) according to the manufacturer’s protocol.

**Lipid Profile Analysis**

Blood samples were collected in EDTA for lipid analysis. HDL was performed using an enzymatic in-vitro assay for the direct quantitative determination of human HDL-cholesterol on Roche automated clinical chemistry analyzers following by the manufacture’s protocol. Triglyceride and cholesterol were performed using VITROS TRIG slides and VITROS chemistry products calibrator kit 2 on VITROS chemistry systems (VITROS 950 Chemistry System). LDL was calculated by the Friedewald Equation. (24)

**Complete Blood Count (CBC) and White Blood Count (WBC) Differentiation Analysis**

Blood cell analysis was performed using COULTER® LH 780 Hematology Analyzer (Beckman Coulter) following the manufacturer’s protocol.

**Statistical Analysis**

All univariate comparisons of the disease state groups used two-sided nonparametric tests or Chi-squared comparisons of proportions, which did not require Gaussian distributions. Most of the measured characteristics, especially the MP, PACs, and soluble protein levels had strongly non-Gaussian distributions. The disease state group comparisons of the subject characteristics did not assume disease state related trends in the responses. These comparisons used the Kruskal-Wallis (KW) test (nonparametric ANOVA comparisons). The analyses of the PC, PAC, MP, and soluble protein levels assumed trends in the responses relative to the disease state. The assessment of the statistical significance of these trends used the Jonckheere-Terpstra (JT) test (nonparametric trend comparisons). The post-hoc comparisons of differences between individual disease state groups (e.g. ES versus LT) used the Wilcoxon two sample test.

The comparisons of the proportions of patients by gender applied the exact Chi-square test of the equality of proportions across the groups. This test compared the observed proportions in the disease state groups against the hypothesis that the proportions were the same for all the disease state groups. The other characteristic proportions (e.g. statin use) were criteria for exclusion from the H group, thus only the ES and LT groups were compared. The analysis of the MP and PAC relationships to the disease state after adjustment for specific covariates were based on logistical regression models with the covariates already included in the models.

On the basis of the assumption of a similar relationship of the disease state differences and the response variabilities, a power calculation was performed with data from Koga et al. (4) using CD144$^+$ EMPs. The sample size of 11 in ES, 22 in LT, and 18 in H, had ~90% power in detecting the relative disease state differences (control versus Diabetes Mellitus) as noted in the aforementioned study.

**RESULTS**

**Subject Design and Characteristics**

On the basis of previous findings that diabetes is a risk factor for cardiovascular disease and duration with diabetes is a further additive component of that risk (23) we recruited ES and LT patients. Subject characteristics are presented in Table 2. A total of 41 subjects (mean age 57 years) were included. Age and gender did not differ between the three groups. As anticipated, the standard marker of blood glucose control, glycated hemoglobin, HbA1c, was significantly altered between all diabetic and healthy individuals, but could not discriminate between ES and LT groups. The lower levels of LDL observed in the diabetic groups was most likely related to statin use (36% in ES, 67% in LT). To investigate this effect in the diabetic groups, we plotted LDL levels in each group separated by statin use, Figure 2A. LT diabetes receiving statins had reduced LDL levels compared to healthy individuals, indicating that without such therapy LDL levels in this cohort would have been substantially higher. The fact that LDL levels were lower in LT patients without statins versus with statins may be due the small sample size of the former ($n = 7$) compared with the latter ($n = 14$). HDL and Triglycerides
did not differ between the three groups. Although LDL levels were controlled in the LT group, systolic blood pressure was higher (P < 0.01) in LT versus H groups. Importantly, as our studies investigated the role of certain cell-derived markers (particularly monocyte MPs), the monocyte and lymphocyte counts were statistically unchanged among the three groups. Red blood cell count was not altered between ES and H groups but decreased significantly in the LT group. Similarly, hemoglobin levels were significantly lower in the LT versus H group (Table 2).

### Measurement and Characterization of Progenitor Cells and Microparticles

Assessment of PCs, PACs, and total MPs along with a range of cell-specific MPs was performed. A general relationship was observed in which PC and PAC levels decreased, and MPs and most MP subtypes increased with either onset or disease duration (Table 3). The exceptions to the above was that PCs increased from H to the ES state, but all fell below H levels in the LT disease state, however this was not statistically significant (P < 0.05). PACs were detected at significantly lower frequencies than PCs and some significant changes were observed. All three PACs were approximately equivalent between H and ES disease state, but CD133* VEGF-R2 dropped significantly from ES to LT subjects. Significant reductions in triple positive PACs, CD133* CD34* VEGF-R2* were only observed between H and LT groups. It is interesting to note that the number of individuals within each group with undetectable levels of triple positive PACs (CD133* CD34* CDVEGF-R2*) by flow cytometry also rose dramatically with onset of disease, (33% in H, 63% in ES, and 68% in LT).

For cell-derived MPs, levels of circulating EMPs and PMPs were altered significantly between H and LT (P = 0.03). CD14* monocyte derived MPs were neither affected by disease state or duration (Table 3). Changes in AnnV+ MPs measured by flow cytometry and nM of PS+ MPs measured by the plate based assay, were statistically equivalent (P = 0.02). However, the flow cytometry method could detect differences with onset of disease, H vs. ES, whereas the plate-based method detected changes with disease duration, ES and LT. The differing resolution between the two assays maybe due to their separate readouts; cytometry (AnnV+ MPs) counts the number of PS+ MPs whereas the plate captures PS+ MPs and quantifies using a prothrombinase assay. Nonetheless, both assays detected differences between the H and LT. These results demonstrate the possibility of employing a more feasible plate based assay to measure MPs in clinical studies than flow cytometry.

A covariate adjustment was also performed to assess if the observed group differences remained significant after the covariates, age, gender, hypertension, and statin use were included in the regression model. As hypertension and statin use were exclusion criteria within the H group, only age and gender could be used in those comparisons. With adjustment for age and gender, AnnV+ MPs was still significant (P = 0.02) between H and ES as...
were EMPs \((P = 0.03)\) and PMPs \((P = 0.01)\) between H and LT. AnnV\(^+\) MPs were borderline significant \((P = 0.053)\) between H and LT. For comparisons between ES and LT, adjusting for age, gender, hypertension, and statin use, no other covariate improves the model after age is included.

### Evaluating the MP/PC Ratio

The measurement of double and triple positive PACs and of cell derived MPs, by flow cytometry, is both technically challenging and expensive. We investigated the value of measuring single positive PCs by flow cytometry and PS\(^+\) MPs by the plate based assay. CD34\(^+\) PCs displayed borderline significant reductions from H to ES \((P = 0.06)\) whilst the measurement of procoagulant MPs by the plate-based assay displayed a significant upward trend with \((P = 0.02)\) Figure 3.

We assessed whether the ratio of MP/PC offered additional information over the investigation of PCs/PACs or MPs alone. Interestingly, the ratio of PS\(^+\) MPs/CD34\(^+\) PC proved a valuable index to distinguish the subject groups \((P = 0.01)\) (Fig. 3) and this change was more significant than any of the other single PC, PAC, or MP subtypes analyzed (Table 3). We also compared PCs, PACs, and MPs to a range of soluble protein markers. The MP/PC ratio was more predictive than a number of often utilized soluble proteins including CRP, ICAM1, and VCAM1 (Table 4). These soluble proteins were quantified via multiplex assays, a cost effective and efficient method often employed in clinical studies. TNF\(_\alpha\), VEGF, IGFBF-1, SDF-1, and PAI-1 although detectable were not significant (data not shown).

Of interest, EPO concentration was higher in the LT group \((P = 0.004)\), (Table 4) and was not due to exogenous recombinant human EPO (rhEPO) or linked to statin use (Fig. 2B). Those within the LT group had significantly depressed red blood cell (RBC) count and hemoglobin and were therefore approaching an anemic state in comparison to both Healthy and ES (Table 2).

### DISCUSSION

To our knowledge, this is the first study showing PACS decrease and MP increase with both onset and duration of Type 2 diabetes. Additionally, we provide
Evidence for the value of assessing the ratio of MPs to PCs (nM of PS\(^+\) MPs/CD34\(^+\) PCs), a measurement that is clinically feasible and more informative than some standard protein markers. PCs and MPs are not byproducts of cardiovascular disease but active components of the disease, and therefore reflect specific disease pathways. For instance, MPs are not only markers of cellular damage but also active agents in promoting endothelial dysfunction (18) and coagulation (10). A reduction in PCs and PACs indicates a loss of vascular reparative ability (6). In demonstrating that levels of both PCs/PACs and MPs correlate with duration of diabetic disease, our results also provide mechanistic insights into the stepwise etiology of diabetes and its contribution to vascular pathology.

The presence of coronary artery disease (CAD) is often asymptomatic in individuals with diabetes (25). Although recent noninvasive studies indicate that CAD can be detected in significant numbers of these individuals, a routine screening approach has not been shown to be clinically useful or cost effective (26). Biomarkers, and particularly cell-derived biomarkers, may prove to be more predictive of cardiovascular events. The search for cell biomarkers of patients at risk for vascular complications is promising but has not become a part of clinical practice because a rapid, easy to perform cell-based assay has not been validated.

A developing and exciting biomarker strategy is the measurement of MPs and assessment of circulating progenitor cells (5,27). MPs are shed by cells as a consequence of cellular injury resulting from activation or death (apoptosis), thus elevation of plasma MPs may be a surrogate marker for vascular cell damage and EMPs are considered markers for endothelial dysfunction (4,16,17). Elevated MPs are found in coronary artery disease or thrombotic microangiopathies, supporting this hypothesis. Further, some evidence exists that the levels of circulating hematopoietic and progenitor cells are reflective of vascular health (27).

In a prospective observational study of diabetic patients referred for angiography, elevated EMP levels, measured using flow cytometry, were predictive for the presence of coronary artery lesions, odds ratio 3.5 (1.8–6.9) (4). Further, it was a more significant independent risk factor than duration of diabetes, lipid levels, or the presence of hypertension. Interestingly, elevated endothelial MPs (CD144\(^+\)) identified a subpopulation of diabetic patients without typical angina symptoms who had atherosclerotic disease on coronary angiography. They also found that endothelial MP levels inversely correlated with coronary blood flow in response to acetylcholine stimulation of endothelium-dependent vasodilatation (4). Similarly, in another study, EMP levels were 2.5 times higher in the presence of high-risk coronary lesions compared with low risk lesions (28). A recent prospective study demonstrated that EMPs in patients at high risk of cardiovascular disease (by virtue of established risk factors) were independent predictors of future events (17).

The etiology of high levels of MPs and low levels of PC and PACs in patients with diabetes is unclear. Microparticles are constitutively released from the surface of cells but can be triggered by cell activation or apoptosis (29). We hypothesize that the high oxidative stress induced by the diabetic state with enhanced reactive oxygen species (ROS) generation and lipid peroxidation promotes MP release. Endothelial nitric oxide synthase (eNOS) decoupling, leading to a switch from the NO producing enzyme to superoxide production may participate in the production of MP (30,31). In fact, endothelial MPs themselves promote endothelial dysfunction through reduction in NO (13) and stimulation of
superoxide from the endothelium. MPs also contain the NAD(P)H oxidase subunit p22phox (32). This intriguing observation suggests that delivery of NAD(P)H subunits to the endothelium by MPs may enhance superoxide production producing a constant cycle of MP production and ROS generation. The administration of an angiotensin II receptor blocker to patients with Type 2 diabetes decreased levels of monocyte and endothelial derived MPs, but not in hypertensive patients without diabetes (33). The administration of statins also reduces MP number in patients with diabetes (20,34).

The diabetic state impairs PC mobilization from the bone marrow and PACs in this disease state display functional impairment, such as reduced proliferation, adhesion, migration, and incorporation, into tubular structures (35,36). The pathological mechanisms underlying reduced circulating PACs in diabetes include bone marrow dysfunction and shortened survival in peripheral blood (6). Indeed, disease-associated MPs may themselves impact negatively on PACs. In-vitro studies indicate that MPs from hypercholesterolemic patients caused a significant PAC loss over MPs harvested from normocholesterolemic controls (22).

Circulating levels of PACs are stimulated by endogenous factors such as proangiogenic cytokines and erythropoietin (37,38). In this study, patients with diabetes had higher levels of erythropoietin than nondiabetics despite lower levels of PACs and hemoglobin. This observation may be due to impaired response of the bone marrow to erythropoietin, driven by enhanced ROS and inflammatory cytokines associated with the diabetic condition. Another study also found increases in EPO levels in microalbuminuric Type 2 diabetes, that was associated with anemic status and also reduced hemoglobin (39). Furthermore, a pathological alteration in the phosphatidylinositol (PI) 3-kinase/protein kinase-B and endothelial nitric oxide synthase eNOS pathways seen in diabetics may be responsible in part to reduced levels of EPCs (40). Again, higher or pathological MPs seen in diabetic patients may play a causative role in reducing eNOS (16).

Our conclusions are based on the assumption that vascular risk increases with duration of diabetic disease as reported in other studies (4,17,22,23).

Further limitations are that the surface markers for PACs are controversial and may not reflect the ability to proliferate or to be incorporated into vessels in vivo. We did however use multiple cells as per our previously validated PAC assay that allows for assessment of many cell combinations (5). The issue of specificity of surface markers for MP subtype identification and the ability to accurately measure MPs by current flow cytometry methods also poses concern. However, our absolute numbers of CD144+ endothelial MPs are in line with those published by other groups (16,41). Although MPs number less than FMPs in our study, the proportion of EMPs to FMPs is slightly higher than that measured in other studies (42).

CONCLUSIONS

The main findings obtained from the experimental study are as follows. First, we found significant alterations in both PCs/PACs and cell derived MPs in Type 2 diabetes and also with duration of disease. Second, we illustrate the potential value of using the ratio of MPs/PCs, which encompasses two biologically relevant markers that impact functionally on disease progression. Third, we show that this ratio may be more informative than many individual standard protein biomarkers commonly used to stratify individuals at heightened cardiovascular risk. From a clinical standpoint, the results from our investigation indicate that a single platform high throughput, multiplexed flow cytometry assay for hematopoietic progenitors and plate-based assay for MPs is a feasible and cost effective method to identify those individuals at highest risk for cardiovascular events.

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LITERATURE CITED


