White blood cell subpopulation changes and prevalence of neutropenia among Arab diabetic patients attending Dasman Diabetes Institute in Kuwait

Fatima Ali1 *, Faisal Alsayegh1, Prem Sharma1, Mohammad Waheedi2, Tania Bayoud2, Faisal Alrefai3

1 Faculty of Medicine, Health Science Center–Kuwait University, Al-Jabriya, Kuwait, 2 Faculty of Pharmacy, Health Science Center–Kuwait University, Al-Jabriya, Kuwait, 3 Dasman Diabetes Institute, Sharq, Kuwait

* fatimakhalilali@hsc.edu.kw

Abstract

Background
The effects of diabetes mellitus on the differential white blood cell count are not widely studied in the Arab populations. The objective of this cross-sectional, retrospective study is to assess the influence of chronic diabetes mellitus on white blood cell counts, absolute neutrophil (ANC) and lymphocyte counts (ALC) as well as the prevalence of benign ethnic neutropenia among Arabs attending the Dasman Diabetes Institute (DDI) in Kuwait.

Methods and findings
1,580 out of 5,200 patients registered in the DDI database qualified for our study. Age, gender, HbA1c and creatinine levels, estimated glomerular filtration rate as well as average WBC, ANC and ALC levels, presence of diabetes-associated complications and anti-diabetic medications were analyzed. Our results showed the mean value of the WBC was $7.6 \pm 1.93 \times 10^9/L$ (95% CI: 2.95–17.15). The mean ANC was $4.3 \times 10^9/L$ (95% CI: 0.97–10.40) and mean ALC was $2.5 \times 10^9/L$ (95% CI: 0.29–10.80). Neutropenia (ANC: $<1.5 \times 10^9/L$) was detected in fifteen patients (0.94%). Six patients (0.4%) fulfilled the definition of lymphopenia (ALC $<1 \times 10^9/L$). Patients with an HbA1c $\geq 7\%$ and those taking at least 3 anti-diabetic medications showed higher values for ANC and ALC. Patients with diabetes-associated neuropathy or nephropathy displayed higher mean ANC values. Our study was limited by overrepresentation of patients over 50 years old compared to those under 50 as well as selection bias given its retrospective nature.

Conclusions
Our study showed that patients with poorly controlled diabetes displayed higher ANC and ALC levels. In addition, patients with DM-associated complications showed higher ANC levels. This finding would suggest that DM exerts a pro-inflammatory influence on differential
Introduction

Diabetes mellitus (DM) is a chronic metabolic disorder characterized by persistently elevated serum blood glucose levels [1]. In the long term, poorly controlled DM is associated with secondary immunodeficiency and frequent infections, thereby contributing to patient’s morbidity and mortality [2–4]. The worldwide prevalence of DM is estimated to be 6.4% [5]. However, certain countries in the Middle East have reported much higher prevalence in their populations. This includes Kuwait (21.1%), Lebanon (20.2%), Qatar (20.2%) and Saudi Arabia (20.0%) [6]. The influence of chronic diabetes on the immune and hematological systems is the subject of ongoing research [7–10]. Data suggests that DM has a wide detrimental effect on complement function as well as both innate and adaptive immunity [2,8,9,11,12]. DM has been shown to exert an adverse effect on polymorphonuclear cells (PMN) in terms of chemotaxis, phagocytic functions and oxidative burst capabilities [2,13]. Chronically uncontrolled DM has also been shown to possibly impair T lymphocyte function, particularly CD4+ cell proliferative responses to protein antigens [14]. One study found a mild but significant neutropenia preceded and accompanied type I diabetes [15]. Another study showed an inverse relationship between white blood cell (WBC) count and insulin tolerance; hence, leukocytosis was associated with the development of DM [16–18]. The effect of DM on overall neutrophil and lymphocyte numbers within Arab populations has not been reported previously in the literature.

Benign ethnic neutropenia (BEN) is a relatively common condition throughout the Middle East, especially among those of African descent [19]. It is a clinical diagnosis based on persistent neutropenia with normal levels of other white blood cell lines, as well as the absence of secondary causes of neutropenia or hematological disorders [20–23]. There are many reports of the prevalence of BEN in Arab populations. However, the prevalence of this disorder in a diabetic Arab population is lacking. The purpose of this retrospective, cross-sectional study is to assess the effects of DM on white blood cell (WBC) subpopulations and to estimate the prevalence of benign ethnic neutropenia in a cohort of Arab patients attending the Dasman Diabetes Institute (DDI) of Kuwait.

Materials and methods

Study design, data source, population and study protocol approval

DDI is a leading research center for diabetes mellitus in Kuwait. It also operates a large outpatient facility dedicated to managing DM and its complications. Data on patients attending the outpatient clinics between June 2006 and February 2015 was retrospectively collected through the DDI electronic healthcare records, a network database which is routinely updated by treating physicians after each patient encounter. The patient laboratory data was retrieved via the Laboratory Information System (LIS). The data extraction for the sole purpose of research was first approved by the International Scientific Advisory Board (ISAB) at DDI and later by the DDI Ethical Review Committee (ERC) (approval number: RA 2014–040). All data was fully anonymized before access by the authors and the requirement for informed consent was waived by both the ISAB at DDI and the local ethics committee before obtaining the data. Investigators underwent the online training course “Protecting Human Research
Participants”, as per National Institute of Health (NIH) requirements. DDI is a research facility and patients who attend provide informed, written consent to have their data used in all future research. Given the retrospective nature of our study, we were not required to obtain permission directly from every patient. Instead, DDI ethics committee granted our group access their database and waived the requirement to obtain informed consent. All data was anonymized and the IRB waived the requirement for informed consent.

Inclusion and exclusion criteria
Inclusion criteria included all patients at least 18 years of age, of Middle Eastern descent, and who carried a diagnosis of either type I or type II DM for at least one year. All selected patients had active files in DDI and had undergone at least two separate assessments within a year from the end of February 2015. Exclusion criteria included patients of non-Middle Eastern descent, those under 18 years, in addition to those suffering from any known hematological, malignant or infectious conditions at the time of data collection. Patients with incomplete records or a single visit were also excluded.

Data collection
The collected data included patient age, gender, current medications and presence of any DM-associated complications as well as laboratory data. For each patient, two most recent WBC values, including 2 absolute neutrophil counts (ANC) and absolute lymphocyte counts (ALC), were collected from the database and the average value for each population calculated. Creatinine, estimated glomerular filtration rate (eGFR) and glycated hemoglobin (HbA1c) were also collected from each patient.

Definitions
Our study defined diabetes mellitus as an HbA1c value ≥ 7%, as defined by the American Diabetes Association [24]. Polypharmacy was defined when patients reported taking at least two or more medications as part of their diabetes management. Normal WBC range was defined as a value between 4–10 x10^9/L [25]. For ANC, the resultant averages were categorized into 4 groups: average ANC ≥ 2 x10^9/L, 1.5–2 x10^9/L, ANC 1–1.5 x10^9/L, and finally <1 x10^9/L. Neutropenia was defined as an ANC < 1.5 [26]. For ALC, the resultant averages were also categorized into 4 groups: ALC ≥ 5 x10^9/L, ALC 3–5 x 10^9/L, ALC 1–3 x 10^9/L, and finally ALC < 1 x10^9/L. Lymphopenia was defined as an ALC < 1 x10^9/L [27]. A creatinine value of ≥ 115 μmol/L and estimated glomerular filtration rate (eGFR) < 90 mL/min/1.73m^2 were defined as consistent with renal impairment [28].

Statistical analysis
The data management, analysis and graphical presentation were performed using the computer software ‘Statistical Package for Social Sciences, SPSS version 24.0’ (IBM Corp, Armonk, NY, USA). The patients’ distribution according to their demographics, clinical features and laboratory findings were presented as number and percentages for each category. The quantitative or continuous variables were first ascertained for normal distribution assumption, applying the Kolmogorov-Smirnov test, and presented as; mean ± standard deviation (SD) and range for normally distributed variables, and median, range, interquartile (IQ) for skewed data. The mean values for laboratory findings were compared applying independent t-test or ANOVA, or non-parametric tests, depending on normal distribution criteria. The two-tailed probability value ‘p’ < 0.05 was considered statistically significant.
Results

Of 5,200 registered patients at DDI, 1,580 (30.4%) met the inclusion criteria. The demographics and laboratory values are shown in Table 1. The mean creatinine and eGFR results are also shown in Table 1. Diabetes-associated complications and their relation to ANC and ALC values are shown in Table 2. Patients’ reported medications and the presence of polypharmacy are displayed in Table 3. The majority (86.1%) of patients were not receiving insulin.

Table 1. Demographic characteristics and laboratory findings, including mean ANC† and mean ALC†, in Arab diabetic patients attending DDI (n = 1580).

| Characteristics | No | % |
|-----------------|----|---|
| Gender          |    |   |
| Male            | 794| 50.3|
| Female          | 786| 49.7|
| Age             |    |   |
| ≤ 50 years      | 497| 31.5|
| > 50 years      | 1083| 68.5|
| Mean Age ± SD (Range) | 52.3 ± 14.1 (18–71) |
| HbA1c           |    |   |
| High (≥ 7.0)    | 1271| 80.4|
| Low (< 7.0)     | 309 | 19.6|
| Mean HbA1c ± SD (Range) | 8.28 ± 1.50 (4.1–14.4) |
| Creatinine      |    |   |
| < 115           | 1408| 89.7|
| ≥ 115           | 162 | 10.3|
| Mean Creatinine ± SD (Range) | 85.4 ± 52.6 (39–1259) |
| Median (Inter-quartile) | 19.7 (7.7–67.8) |
| eGFR            |    |   |
| < 90            | 397 | 77.2|
| ≥ 90            | 117 | 22.8|
| Mean ± SD (Range) | 72.77 ± 25.49 (6.00–160.00) |
| Median (Inter-quartile) | 74.0 (55.0–87.5) |
| ANC             |    |   |
| 0.00–0.99       | 1  | 0.1|
| 1–1.5           | 14 | 0.9|
| 1.5–2           | 47 | 3.0|
| ≥ 2             | 1492| 96.0|
| Mean ± SD (Range) | 4.28 ± 1.45 (0.97–10.40) |
| Median (Inter-quartile) | 4.15 (3.25–5.13) |
| ALC             |    |   |
| 0.00–0.99       | 6  | 0.4|
| 1–3             | 1256| 80.8|
| 3–5             | 285 | 18.3|
| ≥ 5             | 26  | 0.5|
| Mean ± SD (Range) | 2.46 ± 0.74 (0.29–10.80) |
| Median (Inter-quartile) | 2.37 (1.97–2.86) |

*ANC: Absolute neutrophil count
† ALC: Absolute lymphocyte count

https://doi.org/10.1371/journal.pone.0193920.t001
The mean value of the WBC was $7.6 \pm 1.93 \times 10^9$/L ($2.95–17.15$). The mean ANC and ALC values are shown in Table 1. Fifteen patients (0.94%) showed an ANC value $<1.5 \times 10^9$/L, thereby qualifying as neutropenic. Six patients had an ALC value $<1 \times 10^9$/L, which qualified them as lymphopenic. Fig 1 shows a positive correlation between HbA1c and both ANC and ALC. The relationship between renal function (creatinine and eGFR) with ANC and ALC values are shown in Table 4. Table 5 shows an association between medications and ANC and ALC values; in particular, those on three or more medications demonstrated higher ANC and ALC values (Table 6).

**Discussion**

In this retrospective study of Arab diabetic patients, the mean WBC count was $7.6 \pm 1.93 \times 10^9$/L. This result is comparable to another cross-sectional study of 3,772 diabetic Chinese
patients, which found the mean WBC count was $7.2 \pm 1.7 \times 10^9/L$, although our mean is slightly higher ($p<0.001$). Their study also noted an association between higher WBC counts and longer disease duration, higher BMI, poorer glycemic control and lipid profile as well as increased prevalence of diabetes-associated complications [29]. The role of WBC in the pathophysiology and progression of DM is yet to be fully elucidated. WBC can act as an inflammatory marker and evidence suggests that persistently elevated counts bear a direct relationship with increased insulin resistance [16]. In addition, elevated WBC counts, even when within the

Fig 1. Correlation between mean HbA1C values and mean ALC and ANC among Arab diabetic patients attending Dasman Diabetes Institute.

https://doi.org/10.1371/journal.pone.0193920.g001
normal range, may be an independent risk factor in the development of DM-associated micro- and macrovascular complications [29].

Neutrophil and lymphocyte pattern disturbances in diabetic populations are not widely available in the literature, especially with regards to Arab populations where the disease is common. Our study found 0.94% of patients had neutropenia, as defined by a persistent ANC \(< 1.5 \times 10^9/L\). Drug-induced neutropenia was considered a possible underlying cause in those patients. A review of their medications showed some of these patients were taking a sulphonylureas and/or a meglitinide agent for glucose regulation. There are rare associations of WBC disturbances, particularly leukopenia, with these medications (< 0.1% and < 1%, respectively). However, a literature search and review of adverse drug reports revealed no associations between those two medications and neutropenia [30,31]. Therefore, drug-induced neutropenia was considered an unlikely underlying cause in our study cohort. BEN was considered as an alternative diagnosis as these patients demonstrated persistent neutropenia and their records did not indicate the presence of any infection or malignancy, thereby fulfilling the criteria for the diagnosis [19]. BEN is a clinical diagnosis [20, 21]. However, one study found a significant association between persistent neutropenia in patients of African ancestry and polymorphism in the Duffy antigen receptor for chemokines (SNP rs2814778 at chromosome 1q23.2). This study found that the null form of this variant diminished the expression of the receptor on red blood cells, and hypothesized that this variation in turn alters the concentrations of chemokines that control neutrophil production [32]. BEN is a relatively common cause of persistent neutropenia among certain ethnicities, particularly those of African and Arab descent. The population prevalence of BEN among Arabs is estimated to range from 10%–15% [19,33], though one study from Saudi Arabia found the incidence of BEN was as high as 20% [34]. However, our study found the incidence of BEN was much lower than reported. This discrepancy may be attributed to three factors. Firstly, the reported prevalence of BEN in Saudi Arabia was based on a relatively small population study of 100 subjects which contrasts with a larger population reported in our study. Secondly, our findings may reflect the heterogeneity of the local Kuwaiti population, in that it is comprised of more varied, both Middle Eastern and non-Middle Eastern ethnicities, compared to other published reports. The

### Table 4. Differences in mean ANC and mean ALC values and patient demographics among Arab diabetic patients attending DDI.

| Factor | ANC | ALC |
|--------|-----|-----|
|        | N   | Mean ± SD (Range) | p-value | Mean ± SD (Range) | p-value |
| Gender |     |                  |         |                  |         |
| Male   | 780 | 4.25 ± 1.51 (0.97–10.40) | 0.173 | 2.39 ± 0.76 (0.29–10.80) | <0.001 |
| Female | 774 | 4.31 ± 1.39 (1.05–9.13)  |         | 2.54 ± 0.72 (0.51–6.28)  |         |
| Age    |     |                  |         |                  |         |
| ≤ 50 years | 488 | 4.10 ± 1.51 (1.05–10.30) | <0.001 | 2.46 ± 0.75 (0.29–5.97)  | 0.486  |
| > 50 years | 1066 | 4.36 ± 1.42 (0.97–10.40) |         | 2.47 ± 0.74 (0.70–10.80) |         |
| HbA1c  |     |                  |         |                  |         |
| < 7.0 | 300 | 4.03 ± 1.39 (1.05–10.30) | 0.001 | 2.32 ± 0.68 (0.29–5.97)  | <0.001 |
| ≥ 7.0 | 1254 | 4.34 ± 1.46 (0.97–10.40) |         | 2.50 ± 0.75 (0.70–10.80) |         |
| Creatinine |     |                  |         |                  |         |
| < 115 | 1390 | 4.22 ± 1.42 (0.97–10.40) | <0.001 | 2.49 ± 0.74 (0.29–10.80) | <0.001 |
| ≥ 115 | 161 | 4.73 ± 1.65 (2.00–10.40) |         | 2.21 ± 0.70 (0.73–4.16)  |         |
| eGFR   |     |                  |         |                  |         |
| < 90 | 394 | 4.43 ± 1.42 (0.97–10.40) | 0.037  | 2.45 ± 0.69 (0.90–5.45)  | 0.305  |
| ≥ 90 | 115 | 4.12 ± 1.50 (1.10–10.00) |         | 2.54 ± 0.79 (1.05–5.05)  |         |

https://doi.org/10.1371/journal.pone.0193920.t004
close geographical proximity and historical ties between Saudi Arabia and the African continent could mean that their populations share more genetic similarities, and hence a higher prevalence of BEN, compared to the Kuwaiti population, which share greater genetic similarities with those from Asia, Africa and Europe. [35]. Thirdly, Patient selection may be another factor in the low incidence of BEN in our study. Our study exclusively comprised patients with DM, whereas other studies involved healthy populations. Diabetes in this case becomes a confounding factor, in that it establishes a low-grade inflammatory state, characterized by elevated levels of acute phase reactants such as CRP, interleukin-6 (IL-6), secretory phospholipase A2 and tumor necrosis factor alpha (TNFα) [36,37]. These inflammatory agents may subsequently raise neutrophil counts in patients with underlying BEN, thereby masking the true prevalence of this condition.

Table 5. Medications prescribed and mean ANC and mean ALC results among Arab diabetic patients attending DDI.

| Medicine    | N  | Mean ± SD | p-value | Mean ± SD | p-value |
|-------------|----|-----------|---------|-----------|---------|
|             |    | ANC       |         | ALC       |         |
| Aspirin     |    |           |         |           |         |
| Yes         | 515| 4.46 ± 1.51| 0.001  | 2.54 ± 0.81| 0.004  |
| No          | 1039| 4.19 ± 1.42| 2.42 ± 0.70|
| Plavix      |    |           |         |           |         |
| Yes         | 100| 4.42 ± 1.42| 0.310  | 2.37 ± 0.76| 0.227  |
| No          | 1454| 4.27 ± 1.45| 2.47 ± 0.74|
| Warfarin    |    |           |         |           |         |
| Yes         | 4  | 4.15 ± 0.54| 0.669  | 2.76 ± 1.18| 0.645  |
| No          | 1550| 4.28 ± 1.45| 2.46 ± 0.74|
| Metformin   |    |           |         |           |         |
| Yes         | 370| 4.33 ± 1.37| 0.450  | 2.52 ± 0.70| 0.111  |
| No          | 1184| 4.26 ± 1.48| 2.45 ± 0.75|
| Sitagliptin |    |           |         |           |         |
| Yes         | 516| 4.42 ± 1.39| 0.004  | 2.54 ± 0.67| 0.005  |
| No          | 1038| 4.21 ± 1.48| 2.42 ± 0.77|
| Vildagliptin|    |           |         |           |         |
| Yes         | 149| 4.37 ± 1.35| 0.390  | 2.51 ± 0.70| 0.380  |
| No          | 1405| 4.27 ± 1.46| 2.46 ± 0.75|
| Insulin     |    |           |         |           |         |
| Yes         | 218| 4.34 ± 1.55| 0.549  | 2.45 ± 0.74| 0.501  |
| No          | 1336| 4.27 ± 1.44| 2.46 ± 0.74|
| Sulfonylurea|    |           |         |           |         |
| Yes         | 117| 4.46 ± 1.51| 0.155  | 2.61 ± 1.02| 0.114  |
| No          | 1437| 4.26 ± 1.45| 2.45 ± 0.71|
| Repaglinide |    |           |         |           |         |
| Yes         | 87 | 4.52 ± 1.41| 0.097  | 2.68 ± 1.12| 0.060  |
| No          | 1467| 4.26 ± 1.45| 2.45 ± 0.71|
| Pioglitazone|    |           |         |           |         |
| Yes         | 19 | 4.19 ± 1.59| 0.808  | 2.49 ± 0.60| 0.867  |
| No          | 1535| 4.28 ± 1.45| 2.46 ± 0.74|
| Liraglutide |    |           |         |           |         |
| Yes         | 141| 4.43 ± 1.32| 0.152  | 2.63 ± 0.70| 0.004  |
| No          | 1413| 4.26 ± 1.46| 2.44 ± 0.75|

https://doi.org/10.1371/journal.pone.0193920.t005
Our study showed patients with HbA1c ≥ 7% had a small but statistically significantly higher ANC counts compared to those with lower HbA1c. In addition, patients with DM-associated neuropathy, nephropathy and microvascular disease, represented by diabetic foot problems, showed higher ANC counts compared to patients without such complications. Our study also showed that patients on three or more anti-diabetic medications, an indication of difficult-to-control DM, had the highest ANC counts compared to patients requiring less medications. These findings suggests that poorly controlled diabetes exerts a stimulatory influence on ANC numbers. Indeed, while DM exerts a detrimental effect on almost every aspect of the innate immune system, including migratory, phagocytic, oxidative and apoptotic activities, there is also evidence to suggest that DM itself creates a pro-inflammatory milieu [7–10,37–39]. One study found that under both spontaneous and lipopolysaccharide-stimulated conditions, neutrophils from type 2 diabetic patients produced higher amounts of the cytokines interleukin-8 (IL-8), interleukin-1β (IL-1β), TNFα and interleukin-1 receptor antagonist (IL-1RA) compared to healthy controls [38]. In addition, DM has been shown to be associated with elevated CRP levels, NF-κB p65 activity, and soluble adhesion molecules such as ICAM, VCAM and E-selectin [39–41]. It is therefore possible that chronic uncontrolled diabetes creates a pro-inflammatory state causing persistent elevation of neutrophils.

Similar to ANC results, our study showed that elevated HbA1c levels were associated with a small but statistically significant increase in ALC numbers. Patients on three or more anti-diabetic medications were also found to have higher ALC numbers compared to patients on less medications. These findings further suggests that uncontrolled chronic diabetes establishes a pro-inflammatory state. However, our study did not find a significant relationship between DM-associated complications and ALC numbers.

The limitations of our study include an overrepresentation of patients over age 50 years compared to younger patients. In addition, our data does not provide the number of years since patient diagnosis, thus precluding our study’s ability to explore the role of the chronicity of DM on WBC disturbances. Furthermore, given the retrospective nature of this study, selection bias and information bias are two confounding factors which may influence our results.

**Conclusion**

Poorly controlled diabetes mellitus, as reflected by elevated HbA1c values and disease complications, is associated with a small but statistically significant elevation of ANC and ALC values. This elevation may reflect an underlying low-grade pro-inflammatory state established by uncontrolled DM. In addition, our study indicates that the prevalence of neutropenia among the diabetic population is lower than what has been previously reported in other Middle
Eastern countries and Africa. This finding may be due to patient selection and as well population size.

**Author Contributions**

**Conceptualization:** Faisal Alsayegh.

**Data curation:** Mohammad Waheedi, Tania Bayoud, Faisal Alrefai.

**Formal analysis:** Prem Sharma.

**Resources:** Mohammad Waheedi, Tania Bayoud, Faisal Alrefai.

**Writing – original draft:** Fatima Ali.

**Writing – review & editing:** Fatima Ali, Faisal Alsayegh.

**References**

1. Alberti KG, Zimmet PZ. Definition, diagnosis and classification of diabetes mellitus and its complications. Part 1: diagnosis and classification of diabetes mellitus provisional report of a WHO consultation. Diabet Med. 1998; 15(7):539–53. https://doi.org/10.1002/(SICI)1096-9136(19980715):7<539::AID-DIA668>3.0.CO;2-S PMID: 9696993

2. Geerlings SE, Hoepelman AI. Immune dysfunction in patients with diabetes mellitus (DM). FEMS Immunol Med Microbiol. 1999; 26(3–4):259–65. PMID: 10575137

3. Tancredi M, Rosengren A, Svensson A-M, Kosiborod M, Pivodic A, Gudbjörnsdóttir S, et al. Excess Mortality among Persons with Type 2 Diabetes. N Engl J Med. 2015; 373(18):1720–32. https://doi.org/10.1056/NEJMoa1504347 PMID: 26510021

4. Afkarian M, Sachs MC, Kestenbaum B, Hirsch IB, Tuttle KR, Himmelfarb J, et al. Kidney Disease and Increased Mortality Risk in Type 2 Diabetes. J Am Soc Nephrol. 2013; 24(2):302–8. https://doi.org/10.1681/ASN.2012070718 PMID: 23362314

5. Shaw JE, Sicree RA, Zimmet PZ. Global estimates of the prevalence of diabetes for 2010 and 2030. Diabetes Research and Clinical Practice. 2010. p. 4–14. https://doi.org/10.1016/j.diabres.2009.10.007 PMID: 19896746

6. Whiting DR, Guariguata L, Weil C, Shaw J. IDF Diabetes Atlas: Global estimates of the prevalence of diabetes for 2011 and 2030. Diabetes Res Clin Pract. 2011; 94(3):311–21. https://doi.org/10.1016/j.diabres.2011.10.029 PMID: 22079683

7. Sannomiya P, Pereira MAA, Garcia-Leme J. Inhibition of leukocyte chemotaxis by serum factor in diabetes mellitus: Selective depression of cell responses mediated by complement-derived chemotactants. Agents Actions. 1990; 30(3–4):369–76. PMID: 2167002

8. Alba-Loureiro TC, Munhoz CD, Martins JO, Cerchiaro GA, Scavone C, Curi R, et al. Neutrophil function and metabolism in individuals with diabetes mellitus. Brazilian J Med Biol Res. 2007; 40(8):1037–44.

9. Krol E, Ageel R, Banue S, Smogorzewski M, Kumar D, Massry SG. Amlodipine reverses the elevation in [Ca2+]i and the impairment of phagocytosis in PMNLs of NIDDM patients. Kidney Int. 2003; 64(6):2188–95. https://doi.org/10.1046/j.1523-1755.2003.03031.x PMID: 14633142

10. Marhoffer W, Stein M, Schleiniker L, Federlin K. Monitoring of polymorphonuclear leukocyte functions in diabetes mellitus—a comparative study of conventional radiometric function tests and low-light imaging systems. J Biol Chem. 1994; 269:165–70. https://doi.org/10.1002/jbc.26909310 PMID: 7942121

11. Ghosh P, Sahoo R, Vaidya A, Chorev M, Halperin JA. Role of complement and complement regulatory proteins in the complications of diabetes. Endocrine Reviews. 2015. p. 272–88. https://doi.org/10.1210/er.2014-1099 PMID: 25859860

12. Hatanaka E, Monteagudo PT, Marrocos MSM, Campa A. Neutrophils and monocytes as potentially important sources of proinflammatory cytokines in diabetes. Clin Exp Immunol. 2006; 146(3):443–7. https://doi.org/10.1111/j.1365-2249.2006.03229.x PMID: 17100763

13. Peleg AV, Weeraratna T, McCarthy JS, Davis TME. Common infections in diabetes: Pathogenesis, management and relationship to glycaemic control. Diabetes/Endocrinology Research and Reviews. 2007. p. 3–13. https://doi.org/10.1002/dmr.692 PMID: 16960917

14. Eibl N, Spatz M, Fischer GF, Mayr WR, Samstag A, Wolf HM, et al. Impaired primary immune response in type-1 diabetes: results from a controlled vaccination study. Clin Immunol. 2002; 103(3 Pt 1):249–59.
15. Valle A, Giamporcaro GM, Scavini M, Stabilini A, Grogan P, Bianconi E, et al. Reduction of circulating neutrophils precedes and accompanies type 1 diabetes. Diabetes. 2013; 62(6):2072–7. https://doi.org/10.2337/db12-1345 PMID: 23349491

16. Vozarova B, Weyer C, Lindsay RS, Pratley RE, Bogardus C, Tataranni PA. High white blood cell count is associated with a worsening of insulin sensitivity and predicts the development of type 2 diabetes. Diabetes. 2002; 51(2):455–61. PMID: 11812755

17. Schmidt MI, Duncan BB, Sharrett AR, Lindberg G, Savage PJ, Offenbacher S, et al. Markers of inflammation and prediction of diabetes mellitus in adults (Atherosclerosis Risk in Communities study): a cohort study. Lancet (London, England). 1999; 353(9165):1649–52.

18. Twilight G, Afek A, Shamiss A, Derazne E, Tzur D, Gordon B, et al. White blood cells count and incidence of type 2 diabetes in young men. Diabetes Care. 2013; 36(2):276–82. https://doi.org/10.2337/dc11-2298 PMID: 22961572

19. Denic S, Showqi S, Klein C, Takala M, Nagelkerke N, Agarwal MM. Prevalence, phenotype and inheritance of benign neutropenia in Arabs. BMC Blood Disord. 2009; 9:3. https://doi.org/10.1186/1471-2326-9-3 PMID: 19323844

20. Newburger PE. Autoimmune and other acquired neutropenias. Vol. 2016, Hematology. 2016. p. 38–42. https://doi.org/10.1182/asheducation-2016.1.38 PMID: 27913460

21. Walkovich K, Boxer LA. How to Approach Neutropenia in Childhood. Pediatr Rev. 2013; 34(4):173–84. https://doi.org/10.1542/pir.34-4-173 PMID: 23547064

22. Hsieh MM, Tisdale JF, Rodgers GP, Young NS, Trimble EL, Little RF. Neutrophil count in African Americans: Lowering the target cutoff to initiate or resume chemotherapy? Vol. 28, Journal of Clinical Oncology. 2010. p. 1633–7. https://doi.org/10.1200/JCO.2009.24.3881 PMID: 20194862

23. Hay D, Hill M, Littlewood T. Neutropenia in primary care. BMJ. 2014;349.

24. American Diabetes Association. Introduction. Diabetes Care. 2015; 38(Supplement_1):S1–2.

25. Duh S-H. LABORATORY REFERENCE RANGE VALUES. Available from: http://www.stedmansonline.com/webFiles/dict-Stedmans28/APP17.pdf

26. Territo M. Neutropenia—Hematology and Oncology—MSD Manual Professional Edition. Available from: https://www.msdmanuals.com/professional/hematology-and-oncology/leukopenias/neutropenia

27. Territo M. Lymphocytopenia—Hematology and Oncology—MSD Manual Professional Edition. Available from: https://www.msdmanuals.com/professional/hematology-and-oncology/leukopenias/lymphocytopenia

28. Malkina A. Chronic Kidney Disease—Genitourinary Disorders—MSD Manual Professional Edition. Available from: https://www.msdmanuals.com/professional/genitourinary-disorders/chronic-kidney-disease/chronic-kidney-disease

29. Tong PC, Lee KF, So WY, Ng MH, Chan WB, Lo MK, et al. White blood cell count is associated with macro- and microvascular complications in Chinese patients with type 2 diabetes. Diabetes Care. 2004; 27(1):216–22. PMID: 14693992

30. Glipizide XL Side Effects in Detail—Drugs.com. Available from: https://www.drugs.com/sfx/glipizide-xl-side-effects.html

31. Repaglinide Tablets—FDA prescribing information, side effects and uses. Available from: https://www.drugs.com/pro/repaglinide-tablets.html

32. Reich D, Nalls MA, Kao WHL, Akyilbekova EL, Tandon A, Patterson N, et al. Reduced neutrophil count in people of African descent is due to a regulatory variant in the Duffy antigen receptor for chemokines gene. PLoS Genet. 2009; 5(1).

33. Weingarten MA, Potthick-Schwartz EA, Brauner A. The epidemiology of benign leukopenia in Yemenite Jews. Isr J Med Sci. 1993; 29(5):297–9. PMID: 8314691

34. Gari M, Dakhakhni M, Gari A, Alshihri E, Al-Jahdali R, Narasimhan K, et al. Incidence and potential causative factors associated with chronic benign leukopenia in the Kingdom of Saudi Arabia. BMC Proc. 2015; 9(Suppl 2 Selected article from the 2nd International Genomic):S1–6561–9–S2–S1. eCollection 2015.

35. Alsmadi O, Thareja G, Alkafel F, Randhaniy A, Al-Mahdi R, Al-Jahdali R, Al-Sawaf A, et al. Genetic Substructure of Kuwaiti Population Reveals Migration History. PLoS One. 2013; 8(9).

36. Dandona P. Inflammation: The link between insulin resistance, obesity and diabetes. Trends in Immunology. 2004. p. 4–7. PMID: 14698276

37. Pickup JC. Inflammation and activated innate immunity in the pathogenesis of type 2 diabetes. Diabetes Care. 2004; 27(3):813–23. PMID: 14988310

38. Sawant JM. Biochemical changes in polymorphonuclear leukocytes in diabetic patients. J PostgradMed. 1993; 39(4):183–6.
39. Tennenberg SD, Finkenauer R, Dwivedi a. Absence of lipopolysaccharide-induced inhibition of neutrophil apoptosis in patients with diabetes. Arch Surg. 1999; 134(11):1229-33-4.

40. Devaraj S, Dasu MR, Jialal I. Diabetes is a proinflammatory state: a translational perspective. Expert Rev Endocrinol Metab. 2010; 5(1):19–28. https://doi.org/10.1586/eem.09.44 PMID: 20204165

41. Ford ES. The metabolic syndrome and C-reactive protein, fibrinogen, and leukocyte count: Findings from the Third National Health and Nutrition Examination Survey. Atherosclerosis. 2003; 168(2):351–8. PMID: 12801619