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Pain rate and social circumstances rather than cumulative organ damage determine the quality of life in adults with sickle cell disease.

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Due to the significant morbidity associated with sickle cell disease (SCD) sickle cell patients have a reduced quality of life (QoL). Even though pain is considered an important determinant of QoL in sickle cell patients, factors such as organ damage and socioeconomic circumstances may also be important. Therefore we determined the contribution of chronic organ damage and sickle cell related complications to QoL and also analysed the effect of vaso-occlusive crises and socio-economic circumstances on QoL. Consecutive adult sickle cell patients were included. QoL was represented in a physical component scale (PCS) and a mental component scale (MCS) and assessed with SF-36 forms. Higher pain rates were related to lower QoL scores. Both occupation and the level of education were significantly related to PCS while no relation with MCS or pain rate was found. Thirty-five percent patients were unemployed as compared to 6% of the general population and 16% of immigrants without SCD. Neither genotype nor the presence of chronic organ damage were significantly related to QoL. In conclusion, a reduced QoL was mainly determined by pain rate, occupation and educational level.

Chronic organ damage, although a major factor determining life expectancy in SCD, was not a determinant of QoL.

Sickle cell disease (SCD) is a hereditary hemoglobinopathy characterized by recurrent microvascular vaso-occlusion and chronic hemolytic anemia resulting in progressive organ damage and reduced life expectancy. [1;2] Given the chronic nature of SCD and the severity of its complications, patients are likely to have a significantly reduced quality of life (QoL).

Intuitively and historically, pain is widely regarded as the major factor in determining QoL in SCD. However, even though previous studies have indeed demonstrated a reduced QoL of sickle cell patients, this was often not or not only attributable to pain. [2] Factors such as anxiety, depression and socioeconomic circumstances may also contribute to the reduced QoL of sickle cell patients. [3-5]

Sickle cell patients develop organ damage irrespective of their frequency of acute painful events which has been related to significant morbidity and increased mortality and it seems likely that SCD related organ damage negatively impacts QoL of sickle cell patients. [6] However, to our knowledge, the impact of chronic organ damage on QoL in SCD has not been previously assessed. Therefore, we prospectively assessed the QoL in a cohort of consecutive sickle cell patients in whom chronic organ damage, pain rate and the history of sickle cell related complications in the last five years was systematically analyzed. In addition, the impact of occupation and educational level on QoL of these patients was studied.
117 adult patients with sickle cell disease (HbSS, HbSβ⁰, HBSC, HbSβ⁺) were eligible for our study. Twenty-three patients were excluded from analysis because they were not able to fill in the SF-36 questionnaire for different reasons (illiterate persons, mental retardation or repeated no show at their appointments). These excluded patients did not differ from the study population concerning outcome variables (pain, disease severity, education and occupation). QoL was assessed in 51 patients with HbSS, 5 with HbSβ⁰, 11 with HbSβ⁺ and 27 patients with HbSC.

The QoL data of our population are presented in table I. As expected, our patient group scored lower on QoL compared to the healthy Dutch population [9] (data not shown). No correlation between the MCS and PCS was found in the sickle cell population (spearman r=0.14; P>0.20). When comparing male to female sickle cell patients the overall PCS appeared to be significantly higher in males as compared to females (P=0.015). We found no significant difference in the overall MCS between males and females. Older patients scored lower on the PCS in comparison to younger patients (P=0.002) while no significant differences in QoL scores between genotypes was found (data not shown).

The QoL scores in relation to pain rate are depicted in table II. With respect to the MCS, the overall score and all subscales, except for mental health, were significantly lower in patients with the highest pain rate (P=0.021). Although significantly lower scores were found on the role-physical (RF) scale (P=0.005) and the general health (GH) scale (P=0.033) in patients with the highest pain rate the overall PCS was not determined by the number of admissions for painful crises (P=0.110). Also analysis by linear regression confirmed that pain rate is significantly associated with MCS and is less likely to influence the PCS (data not shown).

The prevalence of different forms of organ damage in our study population was: PHT: 24.5%, renal failure: 3.2%, microalbuminuria: 21.3%, retinopathy: 31.2%, iron overload: 9.6%, avascular osteonecrosis: 11.7%, leg ulcers: 6.4%, ACS: 25.5%, priapism: 8.5% and stroke: 4.3%. Except from lower QoL scores (for the PCS) in patients with iron overload (p=0.017), no association between any form of organ damage or sickle cell related complications and QoL scores was found (table III). Moreover, when patients with organ damage and complication were divided according the pathogenesis (hemolysis related complication versus vaso-occlusion/ischemia related complications) no significant differences in QoL was observed as compared to the patients without organ damage at all.

With respect to the relation between occupation and educational level and QoL scores, the PCS was strongly associated with occupation and level of education (p<0.001). The
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overall MCS was not related to occupation or the level of education (respectively p=0.206 and p=0.177). Occupation and the level of education were not different between different genotypes and were not related to pain rates.

Sickle cell disease (SCD) is heterogeneous in its clinical presentation with patients being continuously admitted for the management of disease related complications at one end of the spectrum, and patients rarely requiring medical care at the other. [6] Irrespective of the frequency of acute clinical complications, most patients develop accumulating organ damage throughout their lives as result of chronic hemolytic anemia and ongoing vaso-occlusion. [1]

With respect to QoL, previous studies have reported that sickle cell patients have lower QoL scores as compared to race and age matched controls which were related to pain. [18-20] However, its relation to the presence of chronic organ damage has not been described before.

In the present study organ damage appeared not to be related to any of the QoL scores. If patients were grouped according the pathophysiology of their different forms of organ damage or a history of sickle cell related complication as recently proposed by Kato et al., also no relation with QoL scores was demonstrated. [15] Surprisingly, PHT, which has been related to early mortality in SCD, was not related to QoL in our patients. This might be explained by the fact that our study population consists of patients with mild PHT (regurgitation jet flow velocity < 2.9 m/s) which is generally not related to severe complaints or significantly reduced exercise tolerance.

Factors significantly associated with a reduced QoL were pain rate and social circumstances defined as educational level and occupation. Interestingly, pain rate was only associated with a reduced MCS while the social circumstances of the patients were related to the PCS. These data suggest that frequent painful and unpredictable crises are an important psychological burden rather than a physical burden to sickle cell patients as has been suggested previously. [21] In contrast, social circumstances as reflected by occupation and level of education were mainly related to the PCS, suggesting that only patients in good physical performance are able to finish school and find jobs independently of the frequency of admission for painful crises. This was further supported by the finding that no correlation between the MCS and PCS was found in the sickle cell population, which is in contrast to the findings of QoL assessments in healthy individuals. [9] Similar to previous findings of McClish et al. we did not find a correlation between genotype and QoL scores. [20] Also no difference in QoL scores was observed between patients on hydroxyurea and those who are not on hydroxyurea. This might be explained by the fact that in our study population hydroxyurea was only prescribed to reduce painful crises and the occurrence of acute chest
syndrome. By reducing these complications in patients with an otherwise more severe clinical presentation, hydroxyurea may have improved the QoL. This is in line with a previous prospective study of Ballas et al. who demonstrated an improved QoL with the use of hydroxyurea.[22]

Certain limitations of our study should be taken into account. First, by defining pain rate by the amount of clinical admissions, the conclusions of our study may not be extrapolated for the number of painful crises experienced at home which have been reported to occur frequently in sickle cell patients. [23] However, previous studies have shown that pain (during a painful crises as well as chronic pain) is related to QoL in sickle cell patients. [18;20;22] Second, most forms of organ damage and sickle cell related complications were only present in a relative small group of patients which may have underpowered our study to detect an effect of specific forms of organ damage on QoL. Mild pulmonary hypertension, as one of the most relevant forms of organ damage which has been associated with poor prognosis, was present in 24% of all sickle cell patients and was not related to QoL.

In conclusion, the quality of life scores in consecutive sickle cell patients appear to be determined mainly by pain rate and social circumstances. Despite the contribution of organ damage such as (mild) pulmonary hypertension on prognosis and life expectancy, cumulative organ damage does not seem to be an important determinant of QoL in SCD. In terms of clinical presentation, pain rate is the most important factor for the QoL of adult sickle cell patients.

Methods

Adult sickle cell patients visiting the Department of Hematology of the Academic Medical Centre (AMC) in Amsterdam were eligible for our study. Inclusion criteria were: high performance liquid chromatography (HPLC) confirmed diagnosis of homozygous sickle cell anemia (HbSS), sickle-C disease (HbSC), HbSβ^+^-thalassemia (HbSβ^+^-thal) or HbSβ^0^-thalassemia (HbSβ^0^-thal), age 18 years or older and capable of filling in a questionnaire (Dutch or English). All data were collected during a routine outpatient clinic visit.

QoL was assessed by the use of SF-36 forms. The SF-36 is a short-form health survey which has been proven to be valid and reliable in the black population. [7] It has been previously used to determine QoL in adults with SCD. [8] Briefly, it yields eight different scales (physical functioning, role limitations due to physical problems, role limitations due to emotional problems, social functioning, mental health, vitality, bodily pain and general health perceptions) of functional health and well-being as well as psychometrically-based physical and mental health summary measures. The SF-36 is a generic measure, which is not age,
disease or treatment specific. Accordingly, the SF-36 has proven to be useful in surveys of
general and specific populations, comparing the relative burden of diseases. [9] All patients
were asked to complete this questionnaire during a routine out-patient visit. All scale scores
are linearly converted to a 0 to 100 scale, with higher scores indicating higher levels of
functioning or well being. We analyzed the scores of our study population and compared the
data with the scores of the general healthy Dutch population. [9]

Data on social circumstances represented by occupation and level of education were
routinely gathered during the first routine visit for every patient. With respect to occupation
patients were divided in: employed, unemployed or student. Education level was divided in:
high school or less, vocational education/ community college and tertiary education/university.

Pain rate was defined as the number of admissions for treatment of a vaso-occlusive
crisis from January 2002 until January 2007 and was determined by chart review. [6]
Subsequently three groups were arbitrarily defined: patients without, patients with < 1
admission for painful crises per year and patients with 1 or >1 admission for painful crisis
per year.

Organ damage and sickle cell related complications were assessed by systematic
screening and medical record review and defined as follows: *Pulmonary hypertension (PHT)*:
tricuspid regurgitation jet flow velocity (TRV) equal to or higher than 2.5 m/sec in rest
detected by Doppler echocardiography. PHT was considered absent with no or only trace
TRV. [10] *Renal failure*: an estimated creatinine clearance lower than 100 mL/min
(Cockcroft and Gault). *Microalbuminuria*: urinary creatinine (mmol/L) to urinary albumin
(mg/L) ratio higher than 3.5 for males and higher than 2.5 for females confirmed with 24
hours urine collection with microalbuminuria higher than 30 mg/24 hours. [11] *Retinopathy*:
presence of at least mild non-proliferative retinopathy. [12] *Iron overload*: plasma ferritin
level higher than 1000 µmol/L (on at least three occasions during steady state) and a history
of more than 20 transfused packed cells. [13] *Symptomatic avascular osteonecrosis*: local
pain and reduced function with documented osteonecrosis of the femoral or humeral head (hip
or shoulder X-ray) or a history of surgical intervention for osteonecrosis. *Leg ulcers*: chronic
ulcers of the ankle not otherwise explained. *Acute chest syndrome*: defined as described by
Stuart et al. occurring between January 2002 and January 2007. [14] *Priapism*: spontaneous
painful erection requiring hospital care. *Stroke*: history of stroke confirmed by magnetic
resonance imaging or computerized tomography. We analyzed the relation between organ
damage and QoL both for each form of organ damage or complication separate as well as for
groups. Groups were divided in a group with no organ damage or sickle cell related
complications, a group with organ damage or complications due to hemolysis (leg ulcers, PHT, priapism and stroke) or a group with organ damage related to vaso-occlusion (ACS, retinopathy, avascular osteonecrosis, renal failure and microalbuminuria) as proposed previously. [15]

Hematological and biochemical laboratory parameters were assessed at the same visit at which the patient completed the QoL questionnaire. Fetal hemoglobin percentage (HbF%) was determined by cation-exchange high performance liquid chromatography and α-thalassemia screening was performed with a multiplex PCR assay. [16] If not available we used the results from the latest outpatient visit. [17]

All numbers in the tables are medians with corresponding inter quartile ranges (IQR) unless stated otherwise. Difference in continuous data between groups was tested with the Mann-Whitney test. Difference in categorical data between groups was tested with the Chi-square test. P-values ≤0.05 were considered statistically significant. Multiple linear regression analyses were performed to analyze the interaction between the different scales of the SF-36 questionnaire and pain rate. All missing data were considered missing at random. In the case that one or more questions were not answered in a specific scale of the SF-36 questionnaire, the mean value of the remaining questions on that specific scale was used for the missing items. When half or more of the questions were not answered, the results of the specific scale were discarded. Statistical analysis was performed by using SPSS 12.0.2 (SPSS Inc, Chicago, IL).
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Table I. QoL scores of sickle cell patients

|                     | Male           | Female         | P*   |
|---------------------|----------------|----------------|------|
|                     | SCD N=32       | SCD N=61       |      |
| PCS                 | 42.5 (10.9)    | 36.9 (11.9)    | 0.000|
| Physical Function   | 72.5 (24.2)    | 57.4 (28.0)    | 0.000|
| Role-Physical       | 54.8 (43.5)    | 43.8 (43.6)    | 0.000|
| Bodily Pain         | 60.8 (27.4)    | 50.5 (28.5)    | 0.000|
| General Health      | 48.3 (23.3)    | 41.0 (23.1)    | 0.000|
| MCS                 | 48.4 (12.6)    | 43.9 (12.1)    | 0.023|
| Vitality            | 59.5 (20.8)    | 45.4 (19.3)    | 0.000|
| Social Function     | 73.8 (25.3)    | 62.7 (28.4)    | 0.000|
| Role-Emotional      | 70.0 (41.4)    | 57.2 (45.1)    | 0.035|
| Mental Health       | 74.0 (21.3)    | 63.6 (21.3)    | 0.000|

Mean (SD)

* Mann-Whitney-test
## Table II. QoL in relation to pain rate

|                     | None (n=25)       | >0<1 (n=48)       | >=1 (n=21)       | P*  |
|---------------------|-------------------|-------------------|-----------------|-----|
| PCS                 | 45.3 (34.4-52.1)  | 39.6 (29.5-49.3)  | 33.4 (28.8-39.2)| 0.110|
| Physical Function   | 70.0 (35.0-90.0)  | 70.0 (45.0-90.0)  | 60.0 (39.7-75.0)| 0.390|
| Role-Physical       | 87.5 (0.0-100.0)  | 50.0 (0.0-100.0)  | 0.00 (0.00-50.0)| 0.005|
| Bodily Pain         | 62.0 (36.5-92.0)  | 52.0 (32.0-80.0)  | 41.0 (26.5-61.0)| 0.102|
| General Health      | 52.0 (30.0-83.3)  | 37.0 (25.0-57.0)  | 39.5 (21.3-47.0)| 0.033|
| PCS                 | 50.6 (39.3-57.6)  | 50.8 (38.3-57.1)  | 36.7 (30.4-48.2)| 0.021|
| Vitality            | 55.0 (37.5-75.0)  | 50.0 (43.8-65.4)  | 40.0 (31.3-48.8)| 0.022|
| Social Function     | 75.0 (56.3-100.0) | 75.0 (50.0-100.0) | 50.0 (37.5-75.0)| 0.003|
| Role-Emotional      | 100.0 (8.3-100.0) | 100.0 (33.3-100.0)| 0.00 (0.00-100.0)| 0.032|
| Mental Health       | 80.0 (46.0-92.0)  | 72.0 (48.0-89.0)  | 56.0 (32.0-68.0)| 0.319|

Median (IQR)

*Kruskal Wallis Test*
Table III. QoL in relation to organ damage and sickle cell related complications.

| Manifestations       | N    | Present | Absent | P*   | Present | Absent | P*   |
|----------------------|------|---------|--------|------|---------|--------|------|
| Pulmonary hypertension| 23   | 39.8 (11.3) | 38.5 (11.6) | 0.701 | 50.2 (10.3) | 44.3 (12.8) | 0.060 |
| Renal failure        | 3    | 28.4 (14.4) | 39.0 (11.4) | 0.132 | 50.3 (11.1) | 45.2 (12.4) | 0.466 |
| Microalbuminuria     | 20   | 34.9 (10.5) | 39.9 (11.9) | 0.106 | 41.6 (13.6) | 46.6 (12.3) | 0.186 |
| Retinopathy          | 30   | 37.4 (10.6) | 38.6 (11.9) | 0.638 | 47.5 (11.6) | 45.5 (12.4) | 0.463 |
| Iron overload        | 9    | 30.1 (8.3)  | 39.9 (11.8) | 0.017 | 41.4 (12.4) | 45.9 (12.4) | 0.348 |
| Avascular osteonecrosis| 11  | 36.1 (7.7)  | 39.0 (12.1) | 0.444 | 39.8 (13.8) | 46.1 (12.0) | 0.449 |
| Leg ulcers           | 6    | 38.5 (11.9) | 38.6 (11.7) | 0.966 | 40.6 (7.3)  | 45.6 (12.5) | 0.344 |
| Acute chest syndrome | 24   | 35.0 (12.9) | 40.1 (11.2) | 0.057 | 46.3 (10.6) | 45.2 (13.0) | 0.843 |
| Priapism             | 8    | 45.7 (7.8)  | 41.4 (11.8) | 0.425 | 47.4 (11.1) | 48.7 (13.4) | 0.606 |
| Stroke               | 4    | 42.2 (13.0) | 38.7 (11.8) | 0.190 | 45.7 (16.9) | 45.5 (12.3) | 0.429 |

Mean (SD) *

* Mann Whitney U Test