Current status and achievements of Polish haemato-oncology

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Abstract

The number of newly diagnosed haematological malignancies in Polish adults and children is about 9,000 a year, which constitutes about 5.5% of all malignancies in the country. Adult patients with haematological malignancies are diagnosed and treated in 42 institutions in Poland. The scientific and educational support for this activity is provided under the umbrella of the Polish Society of Haematologists and Transfusiologists (PTHIT, Polskie Towarzystwo Hematologów i Transfuzjologów), the Polish Adult Leukemia Group (PALG), the Polish Lymphoma Research Group (PLRG), the Polish Myeloma Study Group (PMSG), the Polish Myeloma Consortium (PMC), and consultants in haematology.
The aim of this position paper is to present the current status and progress in therapy of haematological malignancies in Polish haematology adult centres, focusing on the activity of PALG, PLRG, and PMSG. The achievements of Polish haematology-oncology at the beginning of the third decade of the 21st century are set out in this paper. Polish haematology-oncology today has an important international position based on contributions to the development of knowledge, international cooperation, and a high quality of patient care. In many instances, clinical trials run by Polish collaborative groups have influenced international standards. Polish haematologists have been the authors of treatment recommendations, and their research has indicated areas for further research.

**Key words:** Polish Society of Haematologists and Transfusiologists, Polish Adult Leukemia Group, Polish Lymphoma Research Group, Polish Myeloma Study Group

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**Introduction — haematology in Poland**

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The Polish Society of Haematology (PTH, Polskie Towarzystwo Hematologiczne) was created in 1949 by Prof. Tadeusz Tempka. From the first meeting of the Society in 1950 in Kraków, Prof. Tempka was its President up to 1972. From 1972 to 1976, Prof. Hugon Kowarzyk was the President. He changed the name of the society to the PTHiT in 1975. The next Presidents were: Prof. Józef Japa (1976–1987), Prof. Janusz Hansz (1987–1995), Prof. Wiesław W. Jędrzejczak (1995–2003), Prof. Andrzej Hellmann (2003–2011), Prof. Tadeusz Robak (2011–2019) and Prof. Iwona Hus (since 2019). The Society has biannual meetings. PTHiT meetings were held in Białystok 2001; Gdańsk 2003; Katowice/Wisła 2005; Warsaw 2007; Wrocław 2009; Lublin 2011; Poznań 2013; Szczecin 2015, Warsaw 2017, and Łódź in 2019. The 29th Meeting will be held in Bydgoszcz in 2021.

The position of national consultant in haematology was held consecutively by Prof. Jerzy Holowiecki (1997–2001), Prof. Jerzy Holowiecki (2001–2002), Prof. Wiesław Jędrzejczak (2002–2014), Prof. Dariusz Wołowiec (2014–2016), again Prof. Jędrzejczak (2016–2018), and Prof. Ewa Lech-Marafida (since 2018). The current team of regional consultants comprises: Lidia Usnarska-Zubkiewicz (woj. dolnośląskie), Małgorzata Całbecka (woj. kujawsko-pomorskie), Marek Hus (woj. lubelskie), Katarzyna Brzezińska-Wojciechowska (woj. lubelskie), Marcel Pasiarski (woj. warmińsko-mazurskie), Lidia Gil (woj. wielkopolskie), and Barbara Zdziarska (woj. zachodniopomorskie).

The Polish Adult Leukemia Group was created in 1975 and chaired by Prof. Jerzy Holowiecki up to 2014, followed by Prof. Sebastian Giebel. The Polish Lymphoma Research Group was created in 2008 and chaired by Dr Janusz Meder followed by Prof. Sebastian Giebel. PALG and PLRG include a total of 36 haematology adult centres actively participating in scientific activity.

The aim of this position paper is to present the current status, achievements and progress in the therapy of haematological malignancies in Polish haematology adult centres, focusing on the activity of PALG, PLRG, and PMSG.

**Epidemiology of haematological malignancies in Poland**

**Morbidity**

According to the National Cancer Registry, the number of newly diagnosed malignancies in the Polish population in 2017 was 164,875 including 8,988 (5.45%) haematological malignancies (Table I) [1]. However, it is worth noting that the National Cancer Registry does not include newly diagnosed patients with myelodysplastic syndromes (D46) or myeloproliferative neoplasms such as polycythemia vera (D45), primary myelofibrosis (D47.1), and essential thrombocytopenia (D75.2). Data from the National Health Fund (NFZ, Narodowy Fundusz Zdrowia) from 2014 indicates there were 1,444 new patients with myelodysplastic syndromes and 3,382 patients with myeloproliferative neoplasms [2].

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The National Cancer Registry provides the following general information regarding the epidemiology of the most frequent haematological malignancies [1].
### Table I. Newly diagnosed haematological malignancies and morbidity rates in Poland in 2017 (no age limits) [1]

| ICD-10 | Malignancy                                      | Newly diagnosed (2017) | Morbidity rate | Standardised morbidity ratio |
|--------|------------------------------------------------|------------------------|----------------|-----------------------------|
| C00-D09 | All malignancies                                | 164,875 (100%)         | 429.11         | 323.50                      |
| C81    | Hodgkin’s lymphoma                              | 753 (0.46%)            | 1.96           | 1.90                        |
| C82    | Follicular lymphoma                             | 442 (0.27%)            | 1.15           | 0.93                        |
| C83    | Non-follicular lymphoma                         | 1,535 (0.93%)          | 4.00           | 3.11                        |
| C84    | Other specified types of T/NK-cell lymphoma     | 244 (0.15%)            | 0.64           | 0.51                        |
| C85    | Other and unspecified types of non-Hodgkin’s lymphoma | 848 (0.51%)     | 2.21           | 1.67                        |
| C88    | Malignant immunoproliferative diseases          | 67 (0.04%)             | 0.17           | 0.13                        |
| C90    | Multiple myeloma and malignant plasma cell neoplasms | 1,600 (0.97%)       | 4.16           | 3.02                        |
| C91    | Lymphoid leukemia                               | 1,801 (1.09%)          | 4.69           | 3.78                        |
| C92    | Myeloid leukemia                                | 1,119 (0.68%)          | 2.91           | 2.28                        |
| C93    | Monocytic leukemia                              | 71 (0.04%)             | 0.18           | 0.13                        |
| C94    | Other leukemias of specified cell type          | 375 (0.23%)            | 0.98           | 0.72                        |
| C95    | Leukemia of unspecified cell type               | 57 (0.03%)             | 0.15           | 0.11                        |
| C96    | Other and unspecified malignant neoplasms of lymphoid, haematopoietic and related tissue | 76 (0.05%)       | 0.20           | 0.16                        |
|        | All haematological malignancies (C81–C96)       | 8,988 (5.45%)          | 23.40          | 18.45                       |

The morbidity rate is the frequency or proportion with which a disease appears in a population. Standardised morbidity ratio is the ratio between the observed number of new diagnoses in a study population and the number of new diagnoses that would be expected based on the age- and sex-specific rates in a standard population and the population size of the study population by the same age/sex groups (European population is assumed as standard population); ICD-10 — International Statistical Classification of Diseases and Health-Related Problems; NK — natural killers

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**Leukemia**

Patients with leukemia comprise 2% of malignancies. The number of newly diagnosed patients with leukemia in 2017 was 3,423. There is an age-dependent increasing trend in the number of newly diagnosed patients. The risk of leukemia in children, adolescents and young adults (AYA) is $4/10^5$, and this risk steadily increases after the age of 50.

**Myeloid leukemia**

Myeloid leukemia (acute and chronic, as reported in ICD-10) comprises 0.7% of malignancies. The number of newly diagnosed patients with myeloid leukemia in 2017 was 1,119, with 55% of diagnoses at age 55–79. Patients diagnosed with myeloid leukemia between 2003 and 2005 had a 1-year survival rate of 45.5% in males and 46.6% in females.

**Lymphoid leukemia**

Lymphoid leukemia (acute and chronic, as reported in ICD-10) comprises 1.1% of malignancies. The number of newly diagnosed patients with lymphoid leukemia in 2017 was 1,801. There are two peak incidence periods: childhood and old age. Patients diagnosed with myeloid leukemia between 2003 and 2005 had a 1-year survival rate of 74.7% in males and 77.3% in females.

**Multiple myeloma (MM)**

This comprises 1% of malignancies. The number of newly diagnosed patients with MM in 2017 was 1,600, with 75% of diagnoses in those aged over 60. The peak incidence is for people in their 70s: 20/10^5 in males and 15/10^5 in females. Patients diagnosed with MM between 2003 and 2005 had a 1-year survival rate of 62.9% in males and 66.0% in females.

**Hodgkin’s lymphoma (HL)**

This comprises 0.5% of malignancies. The number of newly diagnosed patients with HL in 2017 was 753. More than 50% of new diagnoses in males, and 65% in females, are made between the ages of 15 and 40. The peak incidences occur between 25–30 and after the age of 70. Patients diagnosed with HL between 2003 and 2005 had a 1-year survival rate of 89.2% in males and 91.5% in females.

**Non-Hodgkin’s lymphoma (NHL)**

This comprises 2% of malignancies. The number of newly diagnosed patients with NHL in 2017 was 3,069. More than 70% of new NHL diagnoses are made after the age of 50. NHL comprises 7.5% in children, and 5% in age 20–44. Patients diagnosed with NHL between 2003 and 2005 had a 1-year survival rate of 68.5% in males and 70.2% in females.

**Acute myeloid leukemia**

**Treatment of newly diagnosed AML**

Therapeutic management in AML depends on prognostic factors, particularly the patient’s age and cytogenetic as
well as molecular features [3]. Achievement of complete remission (CR) after induction treatment is a prerequisite for successful therapy and prolongation of overall survival (OS). In PALG studies, attempts have been made to modify the standard DA-60 induction protocol (daunorubicin [DNR] 60 mg/m² for three consecutive days in combination with cytosine arabinoside [Ara-C] at a dose of 100–200 mg/m²/day for seven days) in order to improve the efficacy of induction treatment in AML patients eligible for intensive chemotherapy.

Two multicentre randomised trials conducted by PALG demonstrated that the addition of cladribine at a dose of 5 mg/m²/day in a two-hour infusion for five consecutive days adhering to the DA induction (DAC protocol) had a beneficial impact in newly diagnosed AML patients ≤60 years of age [3, 4]. In the DAC arm, CR rate was significantly higher (67.5%) and the occurrence of primary resistance to chemotherapy rarer (21%) compared to standard DA-60 therapy (56%; \( p = 0.01 \) and 34%; \( p = 0.004 \), respectively). The addition of cladribine also exerted a significant effect on OS improvement (3-year OS, 45% vs. 33%), particularly in patients >50 years, with high leukocytosis (>50 G/L), and in the group with unfavourable karyotype [4]. A retrospective analysis showed that the DAC treatment was associated with improved CR and OS rates in the subgroup of patients with FLT3-ITD mutation [5]. In AML patients older than 60 years, DAC appeared to be superior in the subgroup aged 60–65 (CR rate: DAC 51% vs. DA 29%; \( p = 0.02 \)). What’s more, patients with good and intermediate comorbidity Index (CCI) were estimated [7]. What's more, patients with good and intermediate comorbidity Index (CCI) were estimated [7].

**Treatment of refractory and relapsed AML**

A significant limitation of AML treatment is the impossibility of achieving CR with standard induction chemotherapy in 20–30% of patients. In more than half of the patients with CR, leukemia relapse can occur within three years of the initial diagnosis.

The results of the multicentre PALG phase II clinical trials have indicated that cladribine and high-dose Ara-C combined with G-CSF (CLAG) and mitoxantrone (CLAG-M) is a highly efficient treatment for refractory and relapsed AML [8, 9]. The results of the subsequent PALG phase II study showed that the CLAG-M protocol has high antileukemic activity and moderate toxicity in poor prognosis patients with primary drug-resistant AML or in those demonstrating early relapse (1CR <6 months) or relapse after stem cell transplantation [10]. Based on these promising results, in the next phase II trial CLAM (cladribine + Ara-C + mitoxantrone ± G-CSF) was used as an early second induction on day 16 based on bone marrow blasts on day 14 in AML patients aged under 60 years who received DAC as first induction. The study results showed that CLAM used as early second induction might improve CR/CRi rates for younger AML patients with poor early response to DAC induction, but this approach may be associated with higher mortality [11].

**Treatment of acute promyelocytic leukemia (APL)**

In Poland, patients with acute promyelocytic leukemia (APL) are treated in cooperation with the PETHEMA (Programa Español para el Tratamiento de las Hemopatías Malignas) group and according to current PETHEMA protocols. A retrospective analysis of a real-life Polish population showed that early death (ED) remains a major problem in APL, and that shortening the time between the initial contact with a healthcare professional and all-trans retinoic acid administration, as well as the use of appropriate supportive care, could improve the outcomes of an unselected APL population [12].

**Currently ongoing clinical trials**

The PALG-AML1/2016 study aims to compare the safety and efficacy of two commonly used induction (DAC vs. DA-90) and salvage (CLAG-M vs. FLAG-IDA) regimens in AML. This trial is also the first international randomised trial regarding AML induction to prospectively evaluate the impact of measurable residual disease (MRD) on overall survival, using multi-modality testing (flow-cytometry, next-generation sequencing, and PCR) of serial samples. (ClinicalTrials.gov Identifier: NCT03257241). In the PALG-AML1-2018 phase I/ib trial, the safety and efficacy of a combination of CPX-351 with cladribine in elderly patients with relapsed/refractory acute myeloid leukemia is being analysed (EudraCT: 2020-002535-29).

**Chronic myeloid leukemia**

In monitoring the treatment results of chronic myeloid leukemia (CML) with the use of tyrosine kinase inhibitors (TKIs), special attention is paid to appropriate follow-up with molecular methods, which should be performed in certified laboratories able to issue the results of real-time quantitative PCR (RQ-PCR) using the international scale (IS). Polish Molecular Laboratories have been organised in the National Network and have joined the European Leukemia Net and EUTOS projects dedicated to the standardisation of quantitative BCR-ABL1 analysis in patients with CML [13]. The National Molecular Reference Laboratory has been established in Kraków, and this has successfully conducted standardisation and certification procedure in
16 Polish laboratories currently issuing results using the IS [14]. Poland was the first European country with registered imatinib generics in 2014. PALG has established a webpage-based registry to evaluate prospectively the efficacy and tolerability of imatinib generics in a large cohort of adult patients. The report after one year of follow-up of 726 patients (99 previously untreated and 627 patients switched from branded to generic imatinib) showed equal efficacy and tolerability of imatinib generics compared to the branded drug [15].

Myeloproliferative neoplasms Ph-negative

The discovery of mutations that finally confirmed the clonality of Ph-negative myeloproliferative neoplasms resulted in the development of molecular studies also in Polish research centres. This allowed for the characterisation of the Polish population in terms of the presence of new mutations [16], as well as the detection of new mutations in atypical exons of the JAK gene and the MPL gene. Also the subject of research interest was the relationship between the occurrence of complications and the diversified course of diseases depending on the changes in the genome.

In 2018, the results of studies on the evolution of the clinical picture of myeloproliferative neoplasms were published, showing no effect of allele burden on the occurrence of polycythemic transformation [17]. As a result of the cooperation of the centres of the Myeloproliferative Neoplasms Working Group, analysis of the incidence of secondary von Willebrand syndrome in myeloproliferative neoplasms has been published, where it has been shown that this haemostatic pathology is not limited to essential thrombocythemia [18]. In 2019, the preliminary results of a multicentre study of Polish patients treated with ruxolitinib due to myelofibrosis were presented.

Myelodysplastic syndromes (MDS)

In Poland, the standard of care of MDS patients adheres to the current ELN recommendations. In 2008, the MDS Working Group within PALG included 960 MDS patients into a retrospective Polish MDS Registry. Since 2009, 2,513 MDS, AML <30% BM blasts and MDS/MPN patients have been registered prospectively. Cytogenetic results availability improved between 2009–2010 and 2018–2019 from 45% to 77%. Serum ferritin (SF) had a significant impact on outcomes. Patients with higher than 1,000 ng/mL SF versus patients with SF <1,000 ng/mL had a median survival of 320 days versus 568 days (p =0.014) [19]. Assuming that azacitidine (AZA) treated patients are at higher risk of serious infection, especially within the first three AZA cycles, between 2009 and 2016 296 patients were retrospectively analysed. It was found that red blood cell transfusion dependency (odds ratio (OR)=2.38), neutropenia <0.8 ×10⁹/L (OR =3.03), platelet count <50 ×10⁹/L (OR =2.63), albumin level <35 g/L (OR =2.04) and ECOG performance status ≥2 (OR =2.19) all had a significant impact on infectious risk. A subset of patients was selected with high risk of infection rate, 73% versus 25%, and worse clinical outcome, 8 versus 29 months survival [20].

Hypereosinophilic syndrome

Hypereosinophilic syndrome (HES) is a group of rare disorders with a unique clinical picture and challenging treatment. Over the last 20 years, we have witnessed an eruption of molecular findings leading to improved understanding of HES pathogenesis. A small proportion of HES patients may have an abnormal T-cell population responsible for the overproduction of eosinophilopoietic cytokines. These patients are defined to have lymphocytic variant HES. Peripheral blood samples from 42 HES patients were studied for the presence of T-cell receptor rearrangement by PCR and aberrant T lymphocytes by flow cytometry. Clonal T-cell rearrangements were detected in 18 individuals (42.8%) whereas an abnormal T-cell population was revealed only in three patients, with the conclusion that T-cell abnormalities are frequently found in HES [21]. Approximately 10% of HES patients present an interstitial deletion in chromosome 4q12 leading to the expression of \( FIP1L1-PDGFR\alpha \) (F/P) — an imatinib-sensitive gene fusion.

Within the Polish Hypereosinophilic Syndrome Study Group, 32 patients were identified as meeting HES criteria and expressing F/P. Male gender greatly predominated (94%) and splenomegaly and pulmonary involvements were most frequently observed. Treatment with imatinib mesylate (IM) resulted in haematological and molecular CR in 100% of the studied patients. The response was rapid and durable. Imatinib at 100 mg weekly was sufficient to maintain CR in long-term follow-up. The updated results after a median of 12 years on IM have confirmed its excellent efficacy and safety. None of the patients exhibited IM resistance or transformed into acute leukemia. Seven patients stopped IM after achieving long-term remission, and two of them remained in CR for more than seven years after IM discontinuation. IM re-initiation leads to second CR in nearly all patients [22, 23].

Acute lymphoblastic leukemia (ALL)

The treatment of ALL in adults has been based on BFM-like protocols in Poland for more than two decades. Since 1996, adult patients in Poland have been treated with a uniform PALG protocol, with treatment outcomes recorded in prospective studies. A randomised trial demonstrated that the use of granulocyte-colony stimulating factor enabled better adherence to chemotherapy and
improved overall survival of ALL patients [24]. In a phase II study, the safety and efficacy of an originally-developed FLAM regimen (fludarabine, cytarabine, and mitoxantrone) was assessed for patients with relapsed/refractory ALL [25]. The PALG 4-2002 study showed that MRD level ≥0.1% of bone marrow cells after induction assessed by flow cytometry should be considered an independent risk factor for treatment decisions in adult ALL [26]. The combination of MRD status with conventional risk stratification system identified a subgroup of patients allocated to the SR group with MRD ≥0.1% after induction who had a risk of relapse of 71% at three years, versus 9% in the remaining subjects (p = 0.001) [26]. The prognostic value of MRD may be further increased when cytogenetic features are included in the model [27]. Consequently, flow cytometric MRD positive status, i.e. ≥0.1% after induction and ≥0.01% after consolidation, is considered an independent risk factor in the current ALL7 PALG protocol which was introduced in October 2018. Since L-asparaginase is one of the core drugs in PALG protocols, recommendations were published on the use of L-asparaginase in ALL [28]. PALG is currently initiating a randomised, multicentre trial to compare obinutuzumab versus rituximab in newly diagnosed CD20-positive B-cell ALL. A Polish group led an international collaboration to establish recommendations regarding the role of allo-HCT in Ph-negative ALL [29].

**Follicular lymphoma**

The treatment of follicular lymphoma (FL) in Poland has been under the supervision of the PLRG since 2008. Two randomised clinical trials (RCT) were conducted, PLRG1 and PLRG4. In the pre-rituximab era, treatment-naive patients with indolent lymphomas, including 28% of patients with follicular lymphoma, were enrolled into phase III PLRG1 RCT, comparing the efficacy of three protocols: cladribine monotherapy, CC (cladribine and cyclophosphamide), and CVP (cyclophosphamide, vincristine, and prednisone). Protocols containing cladribine yielded significantly better overall responses and complete remission rates and progression-free survival (PFS) times, but not OS [34]. PLRG4, phase III RCT, conducted already in the rituximab era, compared two induction chemoimmunotherapy protocols, R-CVP and R-CHOP, followed by rituximab maintenance, for indolent lymphomas [35]. Patients with FL constituted 42%, while patients with marginal zone lymphoma comprised 38%, among 250 enrolled patients. The frequency of response did not differ between the study arms and the time-to-event endpoints i.e. event-free survival (EFS), PFS and OS, were similar. The only indolent lymphoma with a significantly worse outcome was small lymphocytic lymphoma. Five-year EFS for the whole group reached 61% and 56% in the R-CHOP and the R-CVP arms, respectively. The occurrence of early progression of FL (POD24) was similarly frequent in both arms (13.7 vs. 16.7%, p > 0.05). Grade III/IV adverse events occurred more frequently in the R-CHOP arm compared to the R-CVP arm (55.1 vs. 18.2%). Based on the results of this trial, the R-CVP regimen is preferentially used in PLRG centres for follicular lymphoma patients requiring therapy. In earlier years, PLRG centres participated also in studies involving patients with relapsed/refractory indolent lymphomas, including FL, treated either with four weekly doses of rituximab [36] or radioimmunotherapy [37].

**Hodgkin’s lymphoma (HL)**

A new era in the treatment of Hodgkin’s lymphoma (HL) in Poland started in the past decade with the use of positron emission tomography (PET/CT) for staging [38], interim and final response assessment. Centres allied to the PLRG took part in the validation study of the Deauville Scale, conducted a prospective trial assessing the role of very early interim PET (after one cycle) [39], and developed a new chemotherapy regimen composed of bendamustine, gemcitabine and dexamethasone (BGD) [40] for relapsed/refractory HL. BGD is now being assessed in a prospective study (BURGUND, EudraCT: 2017-001966-97) in patients with progressive disease after first line treatment. The PLRG, together with international partners, has proposed a personalised PET-adapted treatment of early non-bulky HL (eHL) using an innovative risk and response-adapted
strategy. The RAFTING study (EudraCT: 2020-002382-33) aims to assess the efficacy of standard treatment followed by nivolumab in high risk eHL patients defined by positive interim PET (iPET) and/or by high (>84 cm³) initial metabolic tumour volume (MTV) and chemotherapy alone in low risk eHL (defined by both a low MTV and a negative iPET) and the rate of HL relapses in low-risk patients that could be salvaged with delayed radiotherapy and nivolumab maintenance.

Diffuse large B-cell lymphoma

Diffuse large B-cell lymphoma (DLBCL) is the most common aggressive B-cell lymphoma subtype, and prognosis depends on the efficacy of first line therapy. Polish Lymphoma Research Group (PLRG) centres have participated in the most important multicentre studies establishing R-CHOP as the present standard of care (Prof. Jan Walewski), and protocols investigating the potential role of Bruton tyrosine kinase inhibitors and immunomodulating agents (Prof. Wojciech Jurczak). PLRG, in a national multicentre study, confirmed the role of R-CHOP in high risk DLBCL patients [41]. Cardiovascular toxicity of doxorubicin was addressed in several studies, confirming premature cardiovascular mortality [42], investigating the role of arterial hypertension [43], and the feasibility of cardioprotection [44]. Pre-existing diabetes was identified as an independent risk factor of adverse prognosis [45]. In a multicentre approach, PLRG investigated the efficacy of the PREBEN regimen (pixantrone, rituximab, etoposide, and bendamustine) in relapsing refractory cases [46].

Hairy cell leukemia

Hairy cell leukemia (HCL), a chronic lymphoproliferative disorder, responds well to treatment, and one course of cladribine (2-chlorodeoxyadenosine, 2-CdA) usually induces a durable CR. However, there are several administration schedules of this drug and no superiority has been shown of one schedule over the others. Robak et al. [47] demonstrated CR obtained in 75% of patients after 5-day intravenous infusions of 2-CdA, and in 76% after 7-day courses. Interruption 2-hour infusions and continuous 24-hour infusions yielded CR in 82.6% and 66.7% of patients respectively. In another study, Robak et al. [48] compared the efficacy and toxicity of a standard 5-day 2-CdA protocol with a schedule of six weekly 2-CdA infusions. Neither the efficacy nor the toxicity profile was significantly different between the groups. In particular, CR was obtained in 76% of patients in the group of daily 2-CdA administration and in 72% in the weekly administration group. Both PFS and OS were similar in both groups. 2-CdA at 0.12 mg/kg in 2-hour i.v. infusion for five days; or alternatively 2-CdA at 0.12 mg/kg in 2-hour intravenous infusion once a week for six weeks are currently considered in Poland as the standard first line treatment for classical HCL.

Peripheral T cell lymphomas

Peripheral T-cell lymphomas (PTCL) are a heterogenous group of rare diseases that are challenging to treat. Patients with PTCL are treated with the CHOP or CHOP-like regimens used as an induction chemotherapy followed by autologous haematopoietic stem cell transplantation (auto-HCT) as a consolidation of first response. To expand the published experience, Czyż et al. [49] conducted a multicentre, retrospective review of 65 patients with PTCL who underwent auto-HCT as a consolidation of first response achieved with either initial induction chemotherapy or salvage chemotherapy. With the median follow-up of 53 months (range 7–157 months), the 5-year OS and PFS for all patients were 61.5% and 59.4%, respectively. Bone marrow involvement at diagnosis and less than partial remission after induction chemotherapy were factors independently predictive for OS and PFS. Maciejka-Klembowska et al. published a data report from the Polish Paediatric Leukemia/Lymphoma Study Group on clinical features and treatment outcomes of PTCL in 10 children [50]. Different regimens, including CHOP and protocols for lymphoblastic lymphoma, were used. The 5-year OS and event-free survival rates were 63.9% and 81%, respectively. Three children underwent allogeneic HCT, and all of them remain alive and in complete remission.

Mantle cell lymphoma

The mantle cell lymphoma (MCL) treatment paradigm has evolved over recent years in Poland. The National Research Institute of Oncology joined the European Mantle Cell Lymphoma Network (EMCLN) and co-performed two landmark academic clinical trials for untreated MCL patients. In the ‘younger’ trial, a comparison of R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone) and autologous haematopoietic cell transplantation (auto-HCT) consolidation versus alternating cycles of R-CHOP and R-DHAP (rituximab, cisplatin, cytarabine, and dexamethasone) with auto-HCT showed improved time-to-treatment failure [median 3.9 vs. 9.1 years and 40% vs. 65% at 5 year (p =0.038)] for the treatment containing cytarabine [51].

The ‘elderly’ trial demonstrated that the recommended treatment in elderly patients is R-CHOP and rituximab maintenance [52, 53]. Median overall survival (OS) after R-CHOP was 6.4 versus 3.9 years after R-FC (rituximab, fludarabine, and cyclophosphamide) (p =0.0054). Patients responding to R-CHOP had median progression-free survival (PFS) and OS of 5.4 and 9.8 years when randomised to rituximab versus 1.9 (p <0.0001) and 7.1 years (p =0.0026) when randomised to interferon maintenance. Based on the
results of this study, rituximab maintenance after immuno-
chemotherapy treatment became the standard procedure in
elderly patients. The Polish Lymphoma Research Group
(PLRG), as the EMLCN sponsor’s delegate in Poland, is cur-
rently conducting two subsequent first line trials: TRIANGLE
for younger and MCL-R2 for elderly patients.

Burkitt’s lymphoma

A major improvement in the outcomes of adult Burkitt’s
lymphoma patients was achieved in Poland after imple-
menting intensive chemotherapy CODOX-M/IVAC within
the UKLG LY06 study [54]. Long-term survival increased
from the previous less than 20% to close to 70%. Toxicity
of treatment was substantial, with the rate of treatment-
related death approaching 10%. Efficacy was further
improved with the introduction of the GMALL-B-ALL/
NHL-2002 protocol of short intensive chemotherapy
combined with rituximab including high-dose methotrex-
ate, cytarabine, and triple intrathecal therapy [55]. This
largest prospective multicentre trial for adult patients with
Burkitt’s lymphoma leukemia involved 363 patients aged
16 to 85 from 98 European centres including the Oncology
Institute in Warsaw. The rate of complete remission was
88%, 5-year survival was 80%, and 5-year progression-
free survival was 71%. There was no lethal toxicity. Given
the high cure rates across a range of prognostic groups
including age, IPI, and feasibility of immunochemotherapy
in elderly patients, the GMALL-B-ALL/NHL-2002 protocol
is now preferentially used in Poland.

Cutaneous lymphomas

The diagnosis and treatment of cutaneous lymphomas
remains a major challenge [56]. Epigenetic dysregulation
seems to play an important role in the development and
progression of Sézary syndrome (SS). The TMEM244 gene
is ectopically expressed in SS patients, SS-derived cell
lines, and, to a lesser extent, in MF. TMEM244 expression
is negatively correlated with the methylation level of its
promoter. TMEM244 expression can be activated in vitro
by the CRISPR-dCas9-induced specific demethylation of
TMEM244 promoter region. Since both TMEM244 expres-
sion and its promoter demethylation can be potentially
used as markers in SS and some other T-cell lymphomas
[57], STAT5 but also STAT6 and to a lesser extent STAT3
seems to be constitutively activated in Cutaneous T-cell
lymphomas (CTCL). Downregulation of STAT5b protein in
advanced-stage CTCL appears to contribute to its patho-
genesis. STATs seem to be a promising target for new
effective therapeutic agents in CTCL [58]. WP1220 is a
synthetic compound that potently inhibits p-STAT3 and
the growth of CTCL cell lines. Topical treatment of index
skin lesions in stage I–III MF have revealed safety and
some efficacy in MF in the Phase 1b study performed at
the Medical University of Gdańsk [59].

Multiple myeloma

The Polish Myeloma Study Group (PMSG) was established
by Prof. Anna Dmoszyńska in cooperation with Prof. Maria
Kraj in 2005. One of the original clinical protocols was the
evaluation of the efficacy and safety of multiple myeloma
(MM) patient therapy with the CTD regimen (cyclophospha-
mide, thalidomide, and dexamethasone) based on a low
dose of thalidomide (100 mg/d) [60]. Other clinical and
multicentre studies of the PMSG were focused on therapy
with lenalidomide, pomalidomide and bortezomib as well
as the identification of novel cyrogenic prognostic fac-
tors [61–66].

The most significant change in the field of multiple my-
eloma in Poland in recent years was the launch of several
prospective clinical trials initiated by Polish investigators.
On the one hand this provided the opportunity to play a sig-
ificant role in the development of new treatment strate-
gies worldwide, and on the other hand it provided access to
non-reimbursed drugs to Polish patients. PMC006 (ATLAS),
a study that the Polish Myeloma Consortium launched in
2017, was the first non-commercial investigator-initiated
clinical study in Poland; it completed the recruitment of its
planned 160 patients in September 2020. This study as-
sessed two different methods of maintenance after auto-
-PBSCT: R versus KRD.

There are three other important studies ongoing: the
PMC007 (OBI1), which assesses the efficacy of obinutu-
zumab in patients with refractory or recurrent Waldenstro-
em macroglobulinemia; the PMC008 (PREDATOR), where
daratumumab is used to treat patients with biochemical
relapse of myeloma; and the PMC010 (COBRA), which as-
sesses the effectiveness of KRD versus RVD in patients with
newly diagnosed multiple myeloma. The Polish Myeloma
Study Group and the Polish Myeloma Consortium are also
active in the development of the MM Registry. Therefore,
the prospective observational study PMC009 (POMOST)
was launched. In this trial, the treatment conditions and
epidemiology of myeloma in Poland are prospectively as-
sessed on a population of 1,500 patients. PMG and PMC
also took part, together with the Łazarski University and the
National Health Fund, in a pilot project to analyse the epi-
demiology and therapy of myeloma based on the sources
of the NFZ [67]. PMSG regularly publishes recommenda-
tions for the diagnosis and treatment of MM. Several re-
respectative nationwide or international studies have been
performed under the auspices of PMSG, including MM pa-
tients with CNS involvement [68], primary and secondary
plasma cell leukemia [69, 70], primary refractory disease
[71], t(14;16) [72], and bicalon MM [73]. As a result, it
has been shown that the outcomes of advanced stage MM

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patients aged 21–40 are comparable to those of patients aged 41–60 [74].

**Autologous haematopoietic cell transplantation**

In 2019, 1,162 auto-HCTs (306 per 10 million inhabitants) were performed in Poland. Throughout the past decade, work has been underway aimed at the introduction and optimisation of a haematopoietic stem cell chemo-mobilisation protocol with the use of intermediate-dose cytarabine (ID-Ara-C) which is currently commonly used in Polish centres. ID-Ara-C refers to: cytarabine as a i.v. infusion at a dose of 0.4 g/m² twice daily on days +1 and +2 (total dose, 1.6 g/m²), with 10 μg/kg filgrastim started on day +5. In a pilot report, its efficacy was shown when given as a second-line salvage mobilisation regimen. In a subsequent retrospective analysis, the greater benefit of ID-Ara-C compared to 4 g/m² cyclophosphamide as first-line mobilisation in patients with myeloma and lymphoma was demonstrated [75]. In a multicentre analysis from the PLRG, the superior efficacy of ID-Ara-C over DHAP plus G-CSF regimen in lymphoma patients was noted [76]. Finally, in a randomised trial, ID-Ara-C was shown to be superior to G-CSF alone for myeloma patients in terms of a greater proportion of patients achieving a CD34⁺ cell yield sufficient for tandem auto-HCT (98% vs. 70%), a higher median number of collected CD34⁺ cells (20.2 vs. 5.9 ×10⁸ cells/kg), and faster haematopoietic recovery after auto-HCT [77].

**Allogeneic haematopoietic cell transplantation**

Six hundred and eighty-six allogeneic hematopoietic stem cell transplantations (allo-HCT) were performed in Poland in 2019 in 17 centres (12 adult and five paediatric). They included 185 transplantations from family members, 423 from unrelated people, and 78 from haploidentical donors. Since 2013, local Polish donors have predominated among unrelated donors (63% in 2019). The Polish unrelated donor registry is one of the biggest in the world, accounting for more than 1.7 million volunteers. The most frequent indication for allo-HCT was acute myeloid leukemia (32% of procedures from family, 36% from unrelated, and 49% from haploidentical), followed by acute lymphoblastic leukemia (21% from family and 20% from unrelated). Poland has been a very active member of the European Society for Blood and Marrow Transplantation (EBMT). In particular, two Polish investigators have been elected chairs of working parties and members of the EBMT’s Scientific Council: Prof. Jan Styczynski (Infectious Diseases Working Party, 2016–2020) and Prof. Grzegorz Basak (Transplant Complications Working Party, 2017–2021). In addition, Prof. Sebastian Giebel since 2006 has been ALL subcommittee chair and secretary of the Acute Leukemia Working Party. Polish authors have substantially contributed to understanding the role of NK cell alloreactivity after allo-HCT [78], genetic polymorphisms contributing to GVHD [79], the role of pharmaco-economics [80], and the management of post-transplant lymphoproliferative disease (PTLD) [81], as well as having developed a number of international recommendations including treatment and outcome of viral infections following allo-HCT [82] and GVHD [83].

**Infectious complications**

Infectious complications constitute the major cause of morbidity and mortality in haematology and transplant patients. Cooperation between the Infection Disease Study Group of PALG with the Polish paediatric group PSPOH and the Infectious Diseases Working Party of EBMT has resulted in a number of publications. Polish epidemiological analysis on stem cell transplant recipients performed by PALG and PTHIT together with the Polish Society of Paediatric Oncology and Haematology (PTOHD, PSPOH) in a nationwide study revealed a high incidence of bacterial infections in children compared to adults (36% vs. 27.6%; p <0.0001), although with a substantial rate of Gram-negative bacteria in adults. Fungal infections were seen also more often in children (25.3% vs. 6.3%; p <0.0001), as well as viral infections (88.0% vs. 74.9%; p <0.0001). Infection-related mortality was lower in children than in adults (7.8% vs. 18.4%; p <0.0001) [84]. Sub-analysis of MM patients confirmed the importance of multidrug bacterial infections (37.5% of Gram-negative; 54% of Gram-positive) during neutropenia after HCT, influencing mortality [85]. Important data regarding mucormycosis in haematology and transplant settings was published recently, based on 10-year observations. Analysis revealed high mortality (82%) in patients undergoing allo-HCT, despite targeted therapy [86]. The treatment of infectious complications in haematology/transplant patients in Poland is based on systematically updated national recommendations. Recent guidance concerns vaccination in adult patients with haematological malignancies and after stem cell transplantation [87–89], the management of invasive fungal infections [90], the management of CMV infections after allo-HCT [91], and the management of infections after CAR T therapy [92].

**Haematopathology**

The National Histopathological Lymphoma Registry project (NHLR) was implemented in Poland in 2014 by haematopathologists from 24 pathology departments in accordance with the 2008 WHO Classification of Tumours of Haematopoietic and Lymphatic Tissues. DLBCL (32.9%; 2,587), CLL/SLL (31.84%; 2,504), HL (13.37; 1,567), PCM (13.32%, 1,561) and MCL (9.04%; 711) were the most frequent...
in a group of 11,718 tested patients. Major differences between NHLR and European and American data on lymphoma subtypes included a higher incidence of DLBCL, and lower incidences of FL and MALT [93]. The next study, based on 2000–2014 data from the Polish National Cancer Registry, confirmed a lower FL incidence rate in Poland compared to other European countries. FL was ranked fourth in incidence (CR 0.72/10⁵, SR 0.87/10⁵) among all reported mature B-cell non-Hodgkin lymphomas, after CLL/ SLL (CR 3.62/10⁵, SR 4.99/10⁵), PCM (CR 3.78/10⁵, SR 4.97/10⁵) and DLBCL, NOS (CR 2.13/10⁵, SR 2.65/10⁵) [94]. A scheme of comprehensive haematopathological diagnostics of aggressive B-cell lymphomas based on morphology, immune profile and evaluation of MYC, BCL2, and BCL6 gene statuses was proposed, and this was introduced in Poland after the publication of revised WHO classifications in 2017 [95].

**Molecular diagnostics**

During the last ten years, progress has been made in the molecular diagnostics of haematological malignancies in Poland in terms of the spectrum of techniques used, as well as its standardisation. In 2010, the standardisation of *BCR-ABL1* measurement by RQ-PCR in CML patients was carried out in eight laboratories in cooperation with the ELN [14]. The results of the *BCR-ABL1* RD mutation analysis in imatinib-resistant patients were also published [96]. The RQ-PCR technique was improved by modification of the e13a2 and e14a2 transcript measurement in 2019 [97]. Similar progress has been made in the Ph-negative MPN. In 2015, the first results of the molecular characteristics of Polish patients were published [16]. The quality of JAK2 V617F quantification was assessed in an international study [98]. A comparison of qPCR and ddPCR sensitivity in the quantification of the *JAK2* V617F mutation allele burden was performed in 2019 [99]. Further progress was made thanks to the introduction of GEP and NGS to myeloid malignancies diagnostics and MRD assessment [100–102]. A development has also been made in the CLL thanks to the cooperation with ERIC (European Research Initiative on CLL) and the creation of a reference laboratories network in Poland.

**Summary**

Polish haemato-oncology in the 2020s has an important international position based on many contributions to the development of knowledge, cooperation, and a high quality of patient care. In many instances, clinical trials run by Polish collaborative groups have influenced international standards. Polish haematologists have been the authors of many treatment recommendations, and their research has suggested areas for further research.

**Authors’ contributions**

SG, JS, LG — design of study. All authors — collection of data and manuscript writing, critical revision and approval.

**Conflict of interest**

None.

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**Ethics**

The work described in this article has been carried out in accordance with The Code of Ethics of the World Medical Association (Declaration of Helsinki) for experiments involving humans; EU Directive 2010/63/EU for animal experiments; Uniform requirements for manuscripts submitted to biomedical journals.

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