Asparaginase-Associated Pancreatitis in Acute Lymphoblastic Leukemia: Results From the NOPHO ALL2008 Treatment of Patients 1-45 Years of Age

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PURPOSE Asparaginase-associated pancreatitis (AAP) is common in patients with acute lymphoblastic leukemia (ALL), but risk differences across age groups both in relation to first-time AAP and after asparaginase reexposure have not been explored.

PATIENTS AND METHODS We prospectively registered AAP (n = 168) during treatment of 2,448 consecutive ALL patients aged 1.0-45.9 years diagnosed from July 2008 to October 2018 and treated according to the Nordic Society of Pediatric Hematology and Oncology (NOPHO) ALL2008 protocol.

RESULTS Compared with patients aged 1.0-9.9 years, adjusted AAP hazard ratios (HRa) were associated with higher age with almost identical HRa (1.6; 95% CI, 1.1 to 2.3; P = .02) for adolescents (10.0-17.9 years) and adults (18.0-45.9 years). The day 280 cumulative incidences of AAP were 7.0% for children (1.0-9.9 years: 95% CI, 5.4 to 8.6), 10.1% for adolescents (10.0 to 17.9 years: 95% CI, 7.0 to 13.3), and 11.0% for adults (18.0-45.9 years: 95% CI, 7.1 to 14.9; P = .03). Adolescents had increased odds of both acute (odds ratio [OR], 5.2; 95% CI, 2.1 to 13.2; P = .0005) and persisting complications (OR, 6.7; 95% CI, 2.4 to 18.4; P = .0002) compared with children (1.0-9.9 years), whereas adults had increased odds of only persisting complications (OR, 4.1; 95% CI, 1.4 to 11.8; P = .01). Fifteen of 34 asparaginase-rechallenged patients developed a second AAP. Asparaginase was truncated in 17/21 patients with AAP who subsequently developed leukemic relapse, but neither AAP nor the asparaginase truncation was associated with increased risk of relapse.

CONCLUSION Older children and adults had similar AAP risk, whereas morbidity was most pronounced among adolescents. Asparaginase re-exposure should be considered only for patients with an anticipated high risk of leukemic relapse, because multiple studies strongly indicate that reduction of asparaginase treatment intensity increases the risk of relapse.

J Clin Oncol 38:145-154. © 2019 by American Society of Clinical Oncology

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Data Supplement

ASSOCIATED

CONTENT

Author affiliations and support information (if applicable) appear at the end of this article.

Accepted on October 10, 2019 and published at ascopubs.org/journal/ jco on November 26, 2019: D0I https://doi. org/10.1200/JC0.19.

INTRODUCTION

Because adult patients now more frequently receive pediatric-inspired acute lymphoblastic leukemia (ALL) treatment, their tolerance to such therapy has become an important issue.^{1,2} Depending on the extent of asparaginase (ASP) exposure, larger trials report asparaginase-associated pancreatitis (AAP) in up to 11% of children with ALL.³⁻¹⁶ Furthermore, premature withdrawal of ASP reduces cure rates,^{34,17} and one of the commonest causes of ASP truncation in children is AAP

because many experience a second AAP after ASP rechallenge.^{3,5,18}

ASP depletes the body of asparagine,¹⁹ interfering with the highly active pancreatic protein synthesis. Acute pancreatitis arises from premature activation of trypsin within pancreatic acinar cells, acinar cell destruction, concomitant local inflammation, and ultimately pancreatic autodigestion.²⁰ However, the direct mechanism behind AAP is unknown, and treatment is mainly supportive.²¹ Although mortality is reported in only a small percentage of patients,^{3,6,7,13,18,22} both



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acute and long-term morbidities after childhood AAP are common. $^{\rm 13,18,22\text{-}24}$

Some clinical^{3-9,15,18,25} and genetic^{5,9,26} risk factors for acute pancreatitis have been proposed, including adolescent age. However, comparative studies of pediatric and adult patients with ALL, as well as studies exploring the AAP-related morbidity and impact on leukemic relapse risk, are missing. We investigated the cumulative incidence, clinical characteristics, and relapse risk in patients with ALL with AAP aged 1.0-45.9 years uniformly treated according to the Nordic Society of Pediatric Hematology and Oncology (NOPHO) ALL2008 protocol.

PATIENTS AND METHODS

TABLE 1. AAP Characteristics

Study Population

A total of 2,448 patients (including 168 patients with AAP) aged 1.0-45.9 years with a diagnosis of either B-cell precursor or T-cell Philadelphia chromosome-negative

ALL between July 2008 and October 2018 were included. All patients were treated according to the NOPHO ALL2008 protocol in Denmark, Estonia, Finland, Iceland, Lithuania, Norway, and Sweden. An inclusion flowchart is presented in the Data Supplement.

All centers complied with mandatory registration of AAP 4 times a year throughout the study period.^{27,28} We identified patients with AAP in the NOPHO ALL registry on April 2, 2019, which provided data on patient, disease, and antileukemic treatment. Detailed questionnaires in relation to the diagnosis, complications, and outcome of AAP were subsequently completed by local clinicians. The data were merged with previously published data from 60% (80/134) of the pediatric cohort.⁵

AAP-Related Definitions

The diagnosis of acute pancreatitis required fulfillment of at least 2 of 3 diagnostic criteria: (1) abdominal pain; (2) serum amylase (total or pancreas specific) or lipase

All	1.0-9.9		
	1.0-5.5	10.0-17.9	18.0-45.0
168	96 (57)	38 (23)	34 (20)
63 (38)	44 (46)	13 (34)	6 (18)
46 (27)	19 (20)	11 (29)	16 (46)
30 (18)	18 (19)	6 (16)	6 (18)
27 (16)	15 (15)	6 (16)	6 (18)
1 (0.5)	0	1 (2.5)	0
1 (0.5)	0	1 (2.5)	0
166 (99)	94 (98)	38 (100)	34 (100)
106 (66)/7	60 (64)/2	28 (80)/3	18 (56)/2
113 (74)/16	59 (68)/9	30 (91)/5	24 (75)/2
36 (26)/27	17 (22)/18	11 (34)/6	8 (26)/3
21 (13)	19 (20)	0	2 (6)
140 (83)	72 (75)	37 (97)	31 (91)
3 (2)	1 (1)	1 (3)	1 (3)
4	4	0	0
34 (20)	28 (29)	3 (8)	3 (9)
15 (44)	11 (39)	2 (67)	2 (67)
	63 (38) 46 (27) 30 (18) 27 (16) 1 (0.5) 1 (0.5) 166 (99) 106 (66)/7 113 (74)/16 36 (26)/27 21 (13) 140 (83) 3 (2) 4 34 (20)	63 (38) 44 (46) 46 (27) 19 (20) 30 (18) 18 (19) 27 (16) 15 (15) 1 (0.5) 0 1 (0.5) 0 1 (0.5) 0 1 (0.5) 0 1 (0.5) 0 1 (0.5) 0 21 (13) 19 (20) 140 (83) 72 (75) 3 (2) 1 (1) 4 4 34 (20) 28 (29)	63 (38) 44 (46) 13 (34) 46 (27) 19 (20) 11 (29) 30 (18) 18 (19) 6 (16) 27 (16) 15 (15) 6 (16) 1 (0.5) 0 1 (2.5) 1 (0.5) 0 1 (2.5) 1 (0.5) 0 1 (2.5) 1 (66 (99) 94 (98) 38 (100) 106 (66)/7 60 (64)/2 28 (80)/3 113 (74)/16 59 (68)/9 30 (91)/5 36 (26)/27 17 (22)/18 11 (34)/6 21 (13) 19 (20) 0 140 (83) 72 (75) 37 (97) 3 (2) 1 (1) 1 (3) 4 4 0 34 (20) 28 (29) 3 (8)

NOTE. All data are No. (%).

Abbreviations: AAP, asparaginase-associated pancreatitis; ASP, asparaginase; HR, high risk; IR, intermediate risk; SR, standard risk.

*Abdominal pain persisted > 72 hours in 80% of the patients with AAP with this information available (124/155, unknown in 13 patients). †Grading included: (1) mild AAP with symptoms and enzyme elevations lasting < 72 hours; (2) severe AAP with symptoms and/or enzyme elevations lasting > 72 hours or hemorrhagic pancreatitis, pancreatic abscess, or pseudocyst; and (3) death from AAP.

[‡]Two patients with planned polyethylene glycol conjugated *Escherichia coli*-derived ASP (PegASP) re-exposure developed a second episode of acute pancreatitis before restart of PegASP (not included). PegASP was replaced with *Erwinia* chrysanthemi-derived ASP due to allergy in 3 patients after the first AAP event, of whom none developed a second AAP. Five patients developed first-time AAP after the last ASP dose in the protocol treatment and, thus, could not be rechallenged with PegASP.

 \geq 3 times the upper normal limit; and (3) ultrasound, computed tomography, or magnetic resonance imaging compatible with pancreatitis according to the international Ponte di Legno consensus criteria.²⁹ The definition of AAP in this study required a diagnosis within 4 weeks after the last ASP injection—the cutoff time point of measurable polyethylene glycol conjugated *Escherichia coli*-derived ASP (PegASP) acitivity.³⁰ Definitions of grading and complications are presented in the Data Supplement and Tables 1 and 2.

ALL2008 Protocol Treatment

Therapy details on the NOPHO ALL2008 protocol have been reported previously.^{2,27,28,31} Treatment was based on stratification into 4 risk groups: standard risk (SR);

intermediate risk (IR); high risk (HR); and high risk with hematopoietic stem cell transplantation (HR-SCT) in first complete remission (CR1), guided by tumor burden at diagnosis, immunophenotype, cytogenetics, CNS involvement, and minimal residual disease levels on treatment days 15, 29, and 79 (or after the second HR block). SR and IR patients received identical PegASP treatment. Of note, children (SR and IR) were randomly assigned to receive PegASP, 1,000 IU/m²/dose intramuscularly, either at 2-week (control arm) or 6-week (experimental arm) intervals from weeks 14 to 33 (ClinicalTrials.gov identifier: NCT00819351). All children received PegASP at 6-week intervals from week 14 after the randomization closed on March 1, 2016.³² The randomization did not influence

Complication	All	1.0-9.9 years	10.0-17.9 years	18.0-45.0 years
SIRS*	103 (72)/24	66 (75)/8	22 (88)/6	15 (63)/10
Body temperature $>$ 38°C or $<$ 36°C/missing	57 (35)/6	35 (37)/2	14 (40)/3	8 (24)/1
Heart rate $>$ 90 beats/min/missing	107 (69)/14	64 (69)/3	24 (71)/4	19 (70)/7
Respiratory rate > 20 breaths/min/missing	43 (37)/53	31 (42)/23	9 (38)/14	3 (17)/16
WBC $>$ 12 \times 10 ⁹ /L or $<$ 4 \times 10 ⁹ /L/missing	98 (61)/7	62 (68)/5	19 (51)/1	17 (52)/1
Systolic blood pressure $\leq 100 \text{ mmHg}$	52 (33)/11	28 (30)/3	17 (49)/3	7 (24)/5
ICU admission	65 (39)	30 (31)	21 (55)	14 (41)
Assisted mechanical ventilation	14 (22)	7 (23)	4 (19)	3 (21)
Vasopressor support/missing	18 (29)/2	6 (20)	7 (37)/2	5 (36)
Need of acute insulin therapy†	32 (19)	11 (11)	14 (37)	7 (21)
Need of permanent insulin therapy	18 (11)	3 (3)	8 (21)	7 (21)
Pancreatic pseudocysts	45 (27)/1	15 (16)/1	17 (45)	13 (38)
Drainage	19 (42)	4 (27)	11 (65)	4 (31)
Recurrent abdominal pain at last follow-up	15 (10)/11	6 (7)/6	7 (23)/4	2 (6)/1
Elevated pancreatic enzymes at last follow-up	9 (6)/19	3 (3)/10	5 (17)/8	1 (3)/1
Imaging compatible with pancreatitis at last follow-up	11 (7)/15	3 (3)/10	6 (18)/4	2 (6)/1
Inflammation/edema‡	3 (27)	2 (67)	0	1 (50)
Pancreatic pseudocysts§	6 (55)	1 (33)	3 (50)	2 (100)
Hemorrhagell	2 (18)	0	2 (33)	0
AAP-related death¶	3 (2)	1 (1)	1 (3)	1 (3)
Any complication#	95 (57)	44 (46)	31 (82)	20 (59)
Any acute complication	90 (54)	40 (42)	30 (79)	20 (59)
Any persisting complication	33 (20)	8 (8)	16 (42)	9 (26)

 TABLE 2.
 AAP-Related Complications

NOTE. All data are No. (%) or No. (%)/missing.

Abbreviations: AAP, asparaginase-associated pancreatitis; ICU, intensive care unit; SIRS, systemic inflammatory response syndrome.

*SIRS definition: \geq 2 of 4 criteria, including body temperature > 38 or < 36°C, heart rate > 90 beats/min, respiratory rate > 20 breaths/min, and WBC > 12 × 10⁹/L or < 4 × 10⁹/L.

†One patient presented with diabetic ketoacidosis at AAP diagnosis.

*Persisting inflammation/edema in 1 patient and development of inflammation/edema after AAP diagnosis in 1 patient.

§Persisting pseudocysts in 3 patients and development of pseudocysts after AAP diagnosis in 3 patients.

IIDevelopment of hemorrhage after AAP diagnosis in 2 patients.

¶AAP diagnosis postmortem in 1 patient.

#Any known complication included acute complications (ICU admission, acute insulin need, development of pancreatic pseudocysts, and AAP-related death) and persisting complications (permanent insulin need, recurrent abdominal pain, elevated pancreatic enzymes at last follow-up, and imaging compatible with pancreatitis at last follow-up).

cure rates.³² Patients aged \geq 18 years and treated at an adult clinic were not included in the PegASP randomization; however, adults from Estonia, Lithuania, and Sweden received PegASP at 6-week intervals from March, September, and April 2016, respectively. The adult centers from the remaining countries continued PegASP treatment according to the control arm. The NOPHO ALL2008 protocol is presented in the Data Supplement.

The study was approved by the regional research ethics committee of the Capital Region of Denmark (Protocol No. H-2-2010-002) and the Danish Data Protection Agency (Journal No. 2012-58-0004). All patients gave written informed consent, and the study was conducted in accordance with the Declaration of Helsinki (version 2008; www.wma.net).

Statistical Analyses

Patients were followed from the time of ALL diagnosis until the date of the first event (relapse, death during induction or in CR1, or second malignant neoplasm [SMN]); SCT in CR1; loss to follow-up/abandonment of therapy; or last follow-up in the registry or April 2, 2019, whichever came first. When exploring AAP, patients were followed from day 30 (the time of the first ASP dose) until the date of the first event including censoring 4 weeks after the last planned ASP injection or 4 weeks after the ASP truncation date, if available, respectively. The reversed Kaplan-Meier method was used to estimate the follow-up time. Cumulative incidences were estimated using the Aalen-Johansen estimator considering relapse, death, and SMN as competing events; the estimates were compared using Gray's test. The body mass index z-scores were calculated accounting for age and sex according to Danish references.³³ Time to first AAP was analyzed in a Cox proportional hazards regression model including relevant preselected clinical characteristics. As a sensitivity analysis of the predefined age groups (1.0-9.9 years, 10.0-17.9 years, and 18.0-45.9 years), new age groups were explored, each containing approximately 25% of the AAP events (1.0-4.9 years, 5.0-8.9 years, 9.0-16.9 years, and 17.0-45.9 years). Investigating potential risk factors of AAP-related complications, preselected clinical variables were included in a multiple logistic regression model of development of any AAP-related complication within \geq 100 days after the AAP diagnosis. To investigate the association between AAP and time to death in CR1 and relapse, we used Cox models with AAP as a timedependent variable and delayed entry on the CR1 date, respectively. As sensitivity analyses, the models were stratified by risk group on day 29 with delayed entry on day 29 or the CR1 date, if later than day 29. In all Cox models, relevant interactions and the proportional hazards assumption were explored. Two-sided P values < .05 were regarded as statistically significant.

RESULTS

Patient and Treatment Characteristics

Following all patients for a median of 245 days (interquartile range [IQR], 186-259), the day 280 cumulative incidence of first-time AAP (168/2,448) was 8.3% (95% CI, 7.0 to 9.9) with all but 1 late AAP included at this time point. The day 280 cumulative incidences were 7.0% (95% CI, 5.4 to 8.6), 10.1% (95% CI, 7.0 to 13.3), and 11.0% (95% CI, 7.1 to 14.9) in patients aged 1.0-9.9 years, 10.0-17.9 years, and 18.0-45.9 years, respectively (P = .03; Data Supplement). However, when analyzing the 4 new age groups, the risk of AAP already rose at 5 years of age, being 5.4% (95% CI, 3.2 to 7.5) in patients aged 1.0-4.9 years, 10.2% (95% CI, 7.2 to 13.1) in patients aged 5.0-8.9 years, 10.4% (95% CI, 7.3 to 13.4) in patients aged 9.0-16.9 years, and 11.3% (95% CI, 7.6 to 14.9) in patients aged 17.0-45.9 years (P < .001; Fig 1). The clinical characteristics of the patients appear in Tables 1 and 3. None of the patients had a prior history of pancreatitis or any known comorbidity, particularly of the liver and pancreas.

AAP occurred within a median of 10 days (IQR, 6-13; range, 0-28) from last PegASP exposure after a median number of 5 PegASP doses in total (IQR, 3-7; range, 1-14). PegASP was replaced with *Erwinia chrysanthemi*-derived ASP because of allergy in 1 patient before the AAP event. The PegASP activity was measurable in 98% of the AAP patients during treatment with this information available (86/88, unknown in 80 patients). Of note, in 6 of the excluded 8 patients with acute pancreatitis occurring more than 4 weeks after the last PegASP dose, acute pancreatitis occurred within 32-90 days (median, 38 days) after the last PegASP dose. The remaining 2 patients developed acute pancreatitis 210 and 589 days, respectively, after the last PegASP dose (Data Supplement).

In a multiple Cox regression analysis (including sex, immunophenotype, and WBC), the hazard of AAP was doubled already since the age of 5 years (Table 4). Hence, patients aged 5.0-8.9 years demonstrated an HRa of 2.3 (95% CI, 1.5-3.6; P < .0001), patients aged 9.0-16.9 years demonstrated an HRa of 2.5 (95% CI, 1.6-3.8, P < .0001), and patients aged 17.0-45.9 years demonstrated an HRa of 2.5 (95% CI, 1.6 to 3.8; P < .0001; Table 4). No difference in the estimates was observed when including the initially excluded 8 patients diagnosed with acute pancreatitis more than 4 weeks after the last PegASP dose as a sensitivity analysis (results not shown). Additionally, the abovementioned associations remained unchanged when stratifying by induction glucocorticoids (prednisolone v dexamethasone) or by day 29 minimal residual disease-guided risk group (SR v IR v HR/HR-SCT), respectively (results not shown).

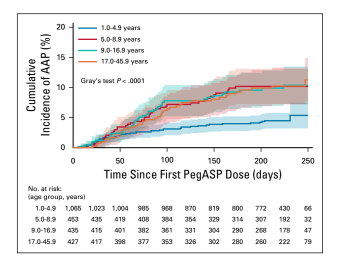


FIG 1. The cumulative incidence of first-time asparaginase-associated pancreatitis (AAP) by age groups with 95% CIs and patients at risk. The day 280 cumulative incidences were 5.4% (95% CI, 3.2 to 7.5) for patients aged 1.0-4.9 years; 10.2% (95% CI, 7.2 to 13.1) for patients aged 5.0-8.9 years; 10.4% (95% CI, 7.3 to 13.4) for patients aged 9.0-16.9 years; and 11.3% (95% CI, 7.6 to 14.9) for patients aged 17.0-45.9 years. Of note, day 280 from acute lymphoblastic leukemia diagnosis to event corresponds to day 250 on the *x*-axis because of entry for all patients on day 30 (the time of the first polyethylene glycol conjugated *Escherichia coli*-derived ASP [PegASP] dose).

AAP-Related Complications and Mortality

Of the 9 patients who presented with hemorrhagic and/or necrotizing pancreatitis at AAP diagnosis, 33% (3/9) had persisting symptoms and signs of chronic pancreatitis at last follow-up, and 1 of these died as a result of AAP (Table 2). Forty-five patients developed pseudocysts (27%; 45/167 with this information available), of whom 21% (9/43; unknown in 2 patients) had recurrent abdominal pain at last follow-up (Table 2).

In a sex-adjusted multiple logistic regression of any AAPrelated complication (ie, AAP-related death; intensive care unit admission; acute and permanent need for insulin therapy; development of pancreatic pseudocysts; recurrent abdominal pain; elevated pancreatic enzymes at last follow-up; and imaging at last follow-up showing pancreatic inflammation/edema, pseudocysts, or hemorrhage), only patients with ≥ 100 days of follow-up after the AAP diagnosis (160/168) were included (median follow-up, 2.3 years; IQR, 1.3-4.2). Patients aged 10.0-17.9 years demonstrated more than 4-fold increased odds of developing any of these AAP-related complications (OR, 4.4; 95% CI, 1.7 to 11.2; P = .002), compared with patients aged 1.0-9.9 years (Table 5). Neither age \geq 18.0 years (OR, 1.5; 95% CI, 0.7 to 3.5; P = .3) compared with children aged 1.0-9.9 years nor sex was associated with development of any AAP-related complication (Table 5). When including the 4 new age groups as a sensitivity analysis, both adolescents aged 9.0-16.9 years and adults aged 17.0-45.9 years had

increased odds of developing any AAP-related complication (9.0-16.9 years: OR, 7.3, 95% Cl, 2.7 to 19.7; P = .0001; and 17.0-45.9 years: OR, 2.6; 95% Cl, 1.1 to 6.4; P = .04; Data Supplement), compared with children aged 1.0-4.9 years.

When analyzing development of any acute AAP-related complication (ie, AAP-related death, acute insulin need, intensive care unit admission, and pancreatic pseudocyst development), only patients aged 10.0-17.9 years had increased odds of developing any acute complication (OR, 5.2; 95% CI, 2.1 to 13.2; P = .0005), compared with patients aged 1.0-9.9 years (Table 5; Data Supplement). Odds of developing any persisting AAP-related complication (ie, elevated pancreatic enzymes at last follow-up; imaging at last follow-up showing pancreatic inflammation/edema, pseudocysts, or hemorrhage; permanent insulin need; and recurrent abdominal pain) were increased for patients aged 10.0-17.9 years (OR, 6.7; 95% CI, 2.4 to 18.4; P = .0002) and patients aged 18.0-45.9 years (OR, 4.1; 95% CI, 1.4 to 11.8; P = .01), compared with patients aged 1.0-9.9 years (Table 5; Data Supplement). Notably, in the sensitivity analysis, patients aged 5.0-8.9 years did not have increased odds of developing any persisting AAP-related complication (OR, 1.7; 95% CI, 0.3 to 10.8; P = .6), compared with patients aged 1.0-4.9 years (Data Supplement).

Death from any cause occurred as the first event in 81 of 2,448 patients, including 5 AAP patients, of whom 3 aged 8.6, 17.3, and 18.6 years died as a result of first-time AAP within 0-29 days from AAP diagnosis (AAP was an unexpected autopsy finding in 1 patient). In an age- and sex-adjusted Cox analysis of time to death in CR1, no difference was found when comparing AAP patients with non-AAP patients (HRa, 0.4; 95% CI, 0.05 to 2.6; P = .3; remaining results not shown). Stratifying the above-mentioned models by risk group on day 29 did not change the results.

Asparaginase Re-Exposure and Relapse

Approximately one fifth (34/168) of the patients with AAP were rechallenged with ASP, of whom 44% (15/34) developed a second episode of AAP after a median of 2 ASP doses (IQR, 1-3; range, 1-7); 40% (6/15) were severe (Table 1). Development of a second AAP episode after ASP re-exposure was not significantly associated with age (Data Supplement). No patient with a second AAP episode was further re-exposed to ASP.

Leukemic relapse in CR1 occurred in 196 of 2,448 patients during the study period, including 21 AAP patients, among whom PegASP was truncated in 81% (17/21) because of AAP. The age- and sex-adjusted hazard of relapse among patients with AAP who were truncated in ASP was not significantly increased when comparing with patients with AAP who were re-exposed

Characteristic	No. (%)	Day 280 Cumulative Incidence of First-Time AAP (%)	95% CI	Р
All patients (delayed entry on day 30)	2,380	8.3	7.0 to 9.9	
Age groups (years; predefined)				.03
1.0-9.9	1,587 (67)	7.0	5.4 to 8.6	
10.0-17.9	415 (17)	10.1	7.0 to 13.3	
18.0-45.9	378 (16)	11.0	7.1 to 14.9	
Age groups (years; sensitivity analysis)				< .001
1.0-4.9	1,065 (45)	5.4	3.2 to 7.5	
5.0-8.9	453 (19)	10.2	7.2 to 13.1	
9.0-16.9	435 (18)	10.4	7.3 to 13.4	
17.0-45.9	427 (18)	11.3	7.6 to 14.9	
Sex				.8
Female	1,019 (43)	7.7	5.9 to 9.4	
Male	1,361 (57)	8.7	6.6 to 10.8	
Immunophenotype				.4
BCP	1,992 (84)	8.8	6.9 to 10.8	
T -cell	388 (16)	6.9	4.2 to 9.6	
WBC				.2
$< 100 \times x10^{9}$ /L	2,056 (86)	8.8	7.0 to 10.7	
$\geq 100 \times x10^{9}/L$	323 (14)	6.3	3.5 to 9.2	
Missing	1			
BMI				.5
≤ -2 SD	117 (5)	7.1	2.4 to 11.8	
> -2 SD to $< +2$ SD	2,066 (87)	8.2	6.6 to 9.8	
\ge +2 SD	164 (7)	9.8	5.1 to 14.6	
Missing	33 (1)			
Induction treatment				.1
Dexamethasone	539 (23)	6.2	4.1 to 8.4	
Prednisolone	1,841 (77)	9.5	7.0 to 12.1	
Treatment group day 29*				.6
SR	1,057 (44)	8.8	5.8 to 11.9	
IR	875 (37)	10.8	5.3 to 16.4	
HR/HR-SCT	429 (18)	7.1	4.5 to 9.8	
Missing	19 (1)	_		

TABLE 3. Baseline Characteristics

Abbreviations: AAP, asparaginase-associated pancreatitis; BCP, B-cell precursor; BMI, body mass index; HR/HR-SCT, high risk/high risk with hematopoietic stem cell transplantation; IR, intermediate risk; SD, standard deviations; SR, standard risk.

*Protocol treatment was modified with the addition of 1 cycle of blinatumomab because of poor treatment response in 1 patient, nelarabine because of poor treatment response in 2 patients, and imatinib because of BCR-ABL translocation in 1 patient.

to ASP (5.0-year cumulative incidence of relapse: 13.2% v 14.2%; HRa, 1.0; 95% Cl, 0.3 to 3.1; P = .97). Moreover, no difference between patients with AAP versus patients without AAP was found in an age- and sex-adjusted Cox analysis (HRa, 1.7; 95% Cl, 0.8 to 3.3; P = .1; remaining results not shown). Stratifying the abovementioned models by risk group on day 29 did not change the results.

DISCUSSION

Despite superior cure rates for children versus adults with ALL, many centers do not use pediatric ALL protocols, partly because of worries about risk of toxicities. Because ASP plays a crucial role in pediatric ALL protocols, the present findings of no increased AAP risk in adults compared with children down to the age of 5 years—in spite of an ASP-heavy protocol—are comforting and in accordance with

TABLE 4. Multiple Cox Regression Analysis of Time to First AAP (n = 2,379) Variable No. AV

Variable	No.	AAP-Specific HRa	95% CI	Р	
Cox model 1					
Age groups (years; predefined)					
1.0-9.9	1,587	ref			
10.0-17.9	415	1.6	1.1 to 2.3	.02	
18.0-45.9	377	1.6	1.1 to 2.4	.02	
Sex					
Female	1,018	ref			
Male	1,361	1.0	0.8 to 1.4	.9	
Immunophenotype					
BCP	1,992	ref			
T cell	387	0.8	0.5 to 1.3	.3	
WBC					
$< 100 \times 10^{9}$ /L	2,056	ref			
$\geq 100 \times 10^{9}$ /L	323	0.8	0.5 to 1.3	.4	
Cox model 2					
Age groups (years; sensitivity analys	is)				
1.0-4.9	1,065	ref			
5.0-8.9	453	2.3	1.5 to 3.6	< .001	
9.0-16.9	435	2.5	1.6 to 3.8	< .001	
17.0-45.9	426	2.5	1.6 to 3.8	< .001	
Sex					
Female	1,018				
Male	1,361	1.0	0.8 to 1.4	.8	
Immunophenotype					
BCP	1,992	ref			
T cell	387	0.7	0.4 to 1.1	.1	
WBC					
$< 100 \times 10^{9}$ /L	2,056	ref			
$\geq 100 \times 10^{9}$ /L	323	0.8	0.5 to 1.3	.4	

NOTE. The table includes 2 Cox models including the predefined age groups and the new age groups as sensitivity analysis, respectively. Abbreviations: AAP, asparaginase-associated pancreatitis; BCP, B-cell precursor; HRa, AAP hazard ratios; ref, reference.

previous studies.^{34,35} Moreover, our findings are compatible with previous pediatric studies reporting more than a 2-fold increased AAP risk in patients older than 9 years of age.^{3,4,8,15,25} This similarity between older children and young adults with ALL has also recently been demonstrated for thromboembolic complications.²

Additionally, the odds of any AAP-related complication were increased by more than 7-fold in adolescents aged 9.0-16.9 years and more than 2-fold in adults aged 17.0-45.9 years, compared with younger children aged 1.0-4.9 years. In fact, adolescents had the most pronounced increase in odds, demonstrating more than 7-fold increased odds of any acute complication and more than 12-fold increased odds of any persisting complication, compared with the youngest children. This emphasizes

the striking vulnerability of this age group, although the reasons hereof are unknown. Changes in endogenous sex hormones may give rise to the increased frequency of insulin resistance and (pre)metabolic syndrome during puberty,³⁶ which has been associated with dyslipidemia and decreased antioxidant capacity.³⁷ Oxidative stress and inflammation play a pivotal role in the pathogenesis of pancreatitis—and probably also in the development of pancreatitis-related complications.^{38,39}

In contrast to age, the role of genetic predisposition is less clear. Recent genome-wide association studies have found different candidate single-nucleotidepolymorphisms associated with pancreatitis in patients with ALL^{5,9,26}; however, only *rs13228878* and *rs10273639* associated with

TABLE 5.	Multiple Logistic Regr	ession of AAP-Re	elated Co	omplications (r	1 = 160
AAP-Rela	ted Complication	No	NR*	95% CI	P

AAP-Related Complication	No.	OR*	95% CI	Р
Any AAP-related complication*				
Age groups (years; predefined)				
1.0-9.9	93	ref		
10.0-17.9	34	4.4	1.7 to 11.2	.002
18.0-45.9	33	1.5	0.7 to 3.5	.3
Sex				
Female	71	ref		
Male	89	1.2	0.6 to 2.3	.6
Any acute AAP-related complication†				
Age groups (years; predefined)				
1.0-9.9	93	ref		
10.0-17.9	34	5.2	2.1 to 13.2	< .001
18.0-45.9	33	1.8	0.8 to 4.1	.1
Sex				
Female	71	ref		
Male	89	1.3	0.7 to 2.6	.4
Any persisting AAP-related complication‡				
Age groups (years; predefined)				
1.0-9.9	93	ref		
10.0-17.9	34	6.7	2.4 to 18.4	< .001
18.0-45.9	33	4.1	1.4 to 11.8	.01
Sex				
Female	71	ref		
Male	89	0.9	0.4 to 2.0	.7

Abbreviations: AAP, asparaginase-associated pancreatitis; OR, odds ratio; ref, reference; ref, reference.

*OR of any AAP-related complication (acute and persisting).

†OR of any AAP-related complication including intensive care unit admission, acute insulin therapy, development of pancreatic pseudocysts, and AAP-related death.

‡OR of any persisting AAP-related complication including permanent insulin therapy, recurrent abdominal pain at last follow-up, elevated pancreatic enzymes at last follow-up, and imaging compatible with pancreatitis at last follow-up.

elevated expression of the *PRSS1* gene encoding for trypsinogen have been validated.²⁶

The early onset of AAP coincided with the PegASP administration, yet the cumulative PegASP dose up to the time of AAP ranged from 1,000 to 14,000 IU/m². Substantial evidence supports that the AAP risk is proportional to the number of doses administered.^{9,32} Notably, this is in contrast to ASP hypersensitivity, which in general occurs after the first few doses.⁴⁰ Although premature withdrawal of ASP has been linked to inferior survival⁴ and PegASP was truncated in the majority of patients with AAP (80%) in our study, neither AAP nor truncation of ASP because of AAP was associated with increased risk of relapse, potentially reflecting low study power in this regard. When looking at ASP re-exposure,

44% of those who were rechallenged with PegASP in our study developed a second event of acute pancreatitis (40% being severe cases), which is in accordance with previous findings in pediatric studies.^{5,18} Thus, current guidelines for children with ALL recommend ASP reexposure only in patients who within 48 hours have no symptoms, normalized pancreatic enzymes, and no evidence of pseudocysts or necrosis.²³ Of note, these guidelines are based on classification of acute pancreatitis according to the original Atlanta criteria-distinguishing between severe (lasting > 48 hours) and nonsevere (lasting \leq 48 hours) acute pancreatitis.^{41,42} For adolescents and adults with ALL, some expert panel guidelines recommend permanent discontinuation of ASP for clinically acute pancreatitis (vomiting and severe abdominal pain with elevated pancreatic enzymes above 3 times the upper normal limit and/or pseudocyst development).⁴³ However, the pressing question in relation to (1) the AAP risk in AAP-naive patients, (2) the risk of a second AAP episode after ASP re-exposure, and (3) overall survival after AAP-related ASP truncation remains unanswered: Who needs more, less, or no ASP? In that respect, the lack of association between characteristics of the first and second AAP is important, which supports that the decision on re-exposure primarily should reflect the anticipated risk of leukemic relapse-except for patients having persisting symptoms from their first AAP.

The main strengths of this study include the international, multicenter, and population-based design and the inclusion of uniformly treated patients with the same diagnosis across a wide age span. Additionally, the online mandatory prospective and systematic registration of 20 predefined treatment-related toxicities strengthens the reliability of the findings.²⁷

The limitations include the lack of power regarding the analyses of leukemic relapse and second AAP event. Moreover, a potential introduction of selection bias in favor of patients with clear-cut symptoms exists because no systematic screening of AAP was performed in patients with abdominal pain or systemic inflammatory response syndrome. Thus, AAP can easily be misinterpreted as sepsis, unless pancreatic enzyme levels are measured.¹⁸ Still, the impact of age on AAP incidence and risk of complications stands strong, and it is unlikely that these potential weaknesses would have markedly influenced the findings.

In conclusion, older children and adults had similar AAP risk, whereas morbidity was most pronounced among adolescents. ASP re-exposure should be considered only for patients with an anticipated high risk of leukemic relapse, because multiple studies strongly indicate that reduction of ASP treatment intensity increases the risk of relapse.^{4,44,45}

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SUPPORT

Supported by research grants from the Research Foundation of Rigshospitalet (University of Copenhagen; CUR), Krista and Viggo Petersen's Foundation (Litra D/6034-29; CUR), the Danish Childhood Cancer Foundation (KS), and the Danish Cancer Society (KS; BOW).

AUTHORS' DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST AND DATA AVAILABILITY STATEMENT

Disclosures provided by the authors and data availability statement (if applicable) are available with this article at DOI https://doi.org/10.1200/JC0.19.02208.

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ACKNOWLEDGMENT

We sincerely thank the acute lymphoblastic leukemia (ALL) patients, colleagues, and research nurses at the ALL centers for contributing to the study, reporting data to the NOPHO ALL registry, and completing the questionnaires. We also thank Kirsten Kørup Rasmussen, Louise Rold Helt, and Pernille Rudebeck Mogensen (Pediatric Research Laboratory, Rigshospitalet, University of Copenhagen, Copenhagen, Denmark) for data extraction help.

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AUTHORS' DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

Asparaginase-Associated Pancreatitis in Acute Lymphoblastic Leukemia: Results From the NOPHO ALL2008 Treatment of Patients 1-45 Years of Age

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Open Payments is a public database containing information reported by companies about payments made to US-licensed physicians (Open Payments).

Cecilie U. Rank Travel, Accommodations, Expenses: Lundbeck

Benjamin O. Wolthers Employment: Novo Nordisk

Birgitte K. Albertsen Honoraria: Shire Research Funding: Erytech Pharma (Inst) Mats M. Heyman Honoraria: Pfizer (Inst), Servier (Inst) Research Funding: Pfizer (Inst), Servier (Inst)

Ulla Wartiovaara-Kautto Honoraria: Pfizer, Sanofi Consulting or Advisory Role: Pfizer, Celgene Travel, Accommodations, Expenses: Roche, Pfizer

No other potential conflicts of interest were reported.