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Glucose levels show independent and dose-dependent association with worsening acute pancreatitis outcomes: Post-hoc analysis of a prospective, international cohort of 2250 acute pancreatitis cases



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# A R T I C L E I N F O

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# ABSTRACT

*Background:* Metabolic risk factors, such as obesity, hypertension, and hyperlipidemia are independent risk factors for the development of various complications in acute pancreatitis (AP). Hypertriglyceridemia dose-dependently elicits pancreatotoxicity and worsens the outcomes of AP. The role of hyperglycemia, as a toxic metabolic factor in the clinical course of AP, has not been examined yet.

*Methods:* We analyzed a prospective, international cohort of 2250 AP patients, examining associations between (1) glycosylated hemoglobin (HbA1c), (2) on-admission glucose, (3) peak in-hospital glucose and clinically important outcomes (mortality, severity, complications, length of hospitalization (LOH), maximal C-reactive protein (CRP)). We conducted a binary logistic regression accounting for age, gender, etiology, diabetes, and our examined variables. Receiver Operating Characteristic Curve (ROC) was applied to detect the diagnostic accuracy of the three variables.

*Results:* Both on-admission and peak serum glucose are independently associated with AP severity and mortality, accounting for age, gender, known diabetes and AP etiology. They show a dose-dependent

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association with severity (p < 0.001 in both), mortality (p < 0.001), LOH (p < 0.001), maximal CRP (p < 0.001), systemic (p < 0.001) and local complications (p < 0.001). Patients with peak glucose >7 mmol/l had a 15 times higher odds for severe AP and a five times higher odds for mortality. We found a trend of increasing HbA1c with increasing LOH (p < 0.001), severity and local complications. *Conclusions:* On-admission and peak in-hospital glucose are independently and dose-dependently

associated with increasing AP severity and mortality. In-hospital laboratory control of glucose and adequate treatment of hyperglycemia are crucial in the management of AP. © 2021 Published by Elsevier B.V. on behalf of IAP and EPC.

#### Introduction

Acute pancreatitis (AP) is the inflammation of the pancreas, most often caused by alcohol consumption and biliary obstruction [1]. The incidence of AP is gradually increasing worldwide, now reported to be 4.6-100/100,000 in the general population [2,3]. The severity of the condition varies, with most cases being classified as mild, but in 15-30% and 10-20% of the cases, moderate and severe disease course will occur, resulting in longer hospitalization, organ failure (OF) and higher mortality (up to 40% in severe cases) [4–6].

Szentesi et al. demonstrated that obesity, hypertension and hyperlipidemia are independent risk factors for several complications in AP. Furthermore, the more components of metabolic syndrome are present, the higher the risk for more severe disease outcomes [7]. A prospective cohort study found that hypertriglyceridemia dose-dependently increases the severity and rate of AP complications [8]. As a next step in filling the gaps of knowledge regarding the role of toxic metabolic factors in the clinical course, we set out to examine the outcome parameters of AP in the context of serum glucose concentration.

Hyperglycemia is an established independent risk factor in numerous diseases. It independently predicts long-term mortality and is associated with a worse prognosis in acute myocardial infarction, irrespective of the presence of diabetes mellitus (DM) [9,10]. It is also associated with a higher rate of poor functional outcomes and less successful revascularization after acute ischemic stroke [11,12]. Furthermore, not only the acute elevations in serum glucose are of interest – studies describe chronic glucose dysregulation to be prognostic for mortality after acute myocardial infarction, both in diabetic and non-diabetic patients [13–15]. In AP, the exact role of glucose dysregulation and its laboratory indicators is yet to be described. However, of the most widely used AP prognostic scoring systems, two (the Ranson and the Glasgow-Imrie) include a serum glucose concentration above 10 mmol/L, highlighting its potential role and association with severity [16].

Our goal was to examine the presence of dose-dependency between glucose dysregulation and clinically important outcomes of AP in a large, multicenter, prospective cohort. In our current cohort analysis (1) previous glucose homeostasis, as assessed by glycosylated hemoglobin (HbA1c) levels; and especially (2) serum glucose levels on admission; and (3) the highest glucose levels during hospitalization have shown dose-dependent associations with key clinically important outcomes of AP. We also established hyperglycemia as an independent and dose-dependent risk factor for mortality.

#### Methods

#### Study design, data source

This study presents a post-hoc analysis of a prospective, international, multicenter registry of AP patients, maintained by the Hungarian Pancreatic Study Group (HPSG). Participants were enrolled in the AP registry if they fulfilled the diagnostic criteria for AP as per the International Association of Pancreatology/American Pancreatic Association (IAP/APA) and HPSG guidelines [17,18]. They were followed up until the end of their AP associated hospital stay (until oral feeding was reinstated without symptoms and with the normalization of laboratory parameters). A list of study sites can be found in the supplementary material (Supplementary Fig. 1, Supplementary Table 1). Between 2012 and 2019, 2461 adult ( $\geq$ 18 years old) AP patients' data were collected. To ensure data quality, a four-tier quality control system was applied, described in detail in a previous publication from the registry [19]. Our study has a partly overlapping population with previous publications from the HPSG AP registry.

### Participants

In the present analysis, we included 2250 patients from the total of 2461 participants in the AP registry with available data on (1) HbA1c any time during the hospitalization with AP and/or (2) on-admission serum glucose measurement and/or (3) at least two serum glucose measurements during hospitalization.

Three variables were taken into account in dividing our examined cohort into subgroups. To observe the role of the glucose homeostasis preceding the admission with AP - 'before AP' (BAP) participants were divided into five groups based on their HbA1c: <6.50; 6.51–7.00; 7.01–8.00; 8.01–9.00; >9.01%. To reflect the onadmission state - 'on-admission AP' (OAP) - seven groups were formed based on on-admission serum glucose levels: <3.99, 4.00-5.99, 6.00-7.79, 7.80-11.09, 11.10-14.99, 15.00-19.99, >20.00 mmol/L. Seven groups were formed based on peak serum glucose during the hospital stay - 'during AP' (DAP) - among those patients who had at least two glucose measurements: <3.99, 4.00-5.99, 6.00-7.79, 7.80-11.09, 11.10-14.99, 15.00-19.99,  $\geq$ 20.00 mmol/L. These boundaries were chosen to reflect already established cut-offs (6.5% and 7% for HbA1c, 4.0 mmol/L, 7.8 mmol/ L and 11.1 mmol/L for glucose, as recommended by the World Health Organization, as well as American and European DM associations/societies [20]), participant and event numbers to maintain statistical power and equal increments to avoid the possibility of arbitrary cut-off selection.

## Representativeness

The distribution of epidemiological data and main outcome variables (such as severity and mortality) were compared between the total registry population and our investigated cohort. There were no significant differences between the total population and the OAP subcohort, indicating that this population's characteristics are representative of the general AP population. We found significant differences from the total cohort: in the length of hospitalization (LOH) in the BAP subcohort, and in age, LOH, and severity in the DAP subcohort (glucose values were probably monitored more closely in more severe cases, and thus often older patients) (Supplementary Fig. 2, Supplementary Table 2).

#### Variables

A complete list of collected variables – including data on laboratory parameters, complications, severity and mortality – is provided in our data quality table in the supplementary material (Supplementary Table 3). Local complications, systemic complications and severity were defined according to the revised Atlanta classification [21].

Glucose measured only within the first 24 hours was accepted as on-admission glucose level. Venous measurements were preferred and accounted for most of the on-admission values, but results from capillary samples were also accepted. HbA1c measured any time during the hospitalization was accepted for this analysis. 95% of measurements occurred within 48 hours of hospitalization. Onadmission values were determined from randomly taken samples, as they usually are, that would mean that they are not fasting samples – however, most AP patients present after hours-days of abdominal pain, nausea and lack of appetite. On the other hand, while most subsequent in-hospital samples are collected early in the morning, a significant proportion of these patients receives enteral feeding. To conclude, not being able to reliably distinguish fasting and non-fasting samples is an inherent limitation of observational trials of AP.

# Statistical analysis

For the descriptive analysis of categorical variables, case number and percentage were computed, while in the case of continuous variables, patient number, mean, standard deviation (SD), median, 25% and 75% quartiles (IQR) were calculated. To identify the three subcohorts' representativeness, we used the Chi-squared test in case of categorical data, the Student's t-test for normally distributed variables, and the Mann-Whitney U test for non-normally distributed variables. In 'Table 1', groups with outlying age (Kruskal-Wallis test) and gender distribution (Fisher exact or Chi-squared test) were sought as well as trends in AP recurrence (RAP), chronic pancreatitis (CP), hyperlipidemia, DM in the medical history (Cochran-Armitage test for trend) and AP episode number among RAP cases (Jonckheere-Terpstra trend test). The dose-dependent effects of HbA1c, on-admission and maximum glucose levels on the investigated outcomes, were tested using the Cochran-Armitage test for trend (in case of categorical variables) and the Jonckheere-Terpstra trend test (in case of continuous variables). Where an overall significant effect was found, a pairwise test of independence (for nominal variables) or a step-down trend test (for continuous variables) with Benjamini-Hochberg p-value corrections was performed as a post-hoc test [22]. Since the overall pvalue was significant in case of age in the OAP subcohort, we performed the Dunn posthoc test to identify the outlying group, applying the above mentioned correction method (Supplementary Table 5).

To detect the predictive accuracy of HbA1c, on-admission glucose level and maximum glucose level on the mortality and severity, the Receiver Operating Characteristic Curve (ROC) was applied. To check the performance of the classification, we used Area Under the Curve (AUC) with 95% confidence intervals (CI) (depending on the AUC value, the accuracy of the test can be categorized as followed: 0.5–0.6 fail, 0.6–0.7 poor, 0.7–0.8 fair, 0.8–0.9 good and above 0.9 excellent). We calculated the potential best cut-off value as well. Odds ratios (OR) with 95% CI were calculated for severe AP cases and mortality. Binary logistic regression was used to test the independent prognostic role of the three investigated variables. The model contains the following parameters in all cases: age, gender, DM, AP etiology and HbA1c, on-admission glucose level or maximum glucose level. We also

created univariate models for severity and mortality with the same parameters, this can be found in our supplementary material (Supplementary Tables 10 and 11). All calculations were performed with the statistical software R, version 4.0.2 (R Core Team, 2020, Vienna, Austria) using the coin (v1.3-4; Hothorn et al., 2008), rcompanion (v2.3.27; Mangiafico, 2021), DescTools (v0.99.39; Signorell et al., 2020), PMCMRplus (v1.9.0; Pohlert, 2021) and pROC (v1.17.0.1; Robin et al., 2011), dunn.test (v1.3.5; Dinno, 2017) packages.

Box plots of LOH and CRP included in the main body of the article do not visualize outliers above a certain value to allow for a better interpretation of the figure. The original figures with all outliers can be found in the supplementary material (Supplementary Fig. 3).

#### Ethical approval

Ethical approval for the registry was granted in 2012 by the Scientific and Research Ethics Committee of the Medical Research Council (22,254–1/2012/EKU). The registry protocol was approved by the institution's human research committee preceding the study initiation. It complies with the ethical guidelines of the Declaration of Helsinki, reaffirmed in 2013. All participants provided written, informed consent for participation.

# Study reporting

This study is reported according to the 'Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) Statement' [23].

# Results

## Study population

A total of 2250 AP cases were analyzed. Table 1 contains the baseline information of participants in each subgroup of all three subcohorts (BAP, OAP and DAP). A statistically significant trend of increasing proportion of hyperlipidemia and DM in the medical history with increasing HbA1c and glucose values was seen in all subcohorts. Among groups with different on-admission glucose values, significant age differences were noted, with the highest mean ages observed in groups 4–6 (glucose 7.8–19.99 mmol/L) and lowest in group 1 (glucose <4 mmol/L).

## Etiology distribution

Fig. 1 shows how different etiologies of AP were distributed across our examined subgroups. While most etiologies showed a balanced distribution, hypertriglyceridemia (HTG) increased in higher HbA1c and serum glucose groups, starting at HbA1c >7% in the BAP, and glucose >11.1 mmol/L in the OAP and DAP subcohorts. A detailed table of etiology distribution can be found in our supplementary material (Supplementary Table 4).

## BAP: pre-existing disturbance of glucose metabolism shows a trend of increasing AP severity and local complications

While no statistically significant differences were noted (p = 0.394), a trend of increasing HbA1c and increasing AP severity was observed. AP severity was highest in Group 4 (21.4% moderate, 5.8% severe). HbA1c was directly associated with the length of hospitalization (p < 0.001) and maximal CRP (p < 0.001), both peaking in group 4 probably due to the higher proportion of moderate cases, but not with mortality, which was the greatest in

#### Table 1

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Baseline characteristics of participants in each group of all three subcohorts. AP: acute pancreatitis; BAP: before AP; OAP: on-admission with AP; DAP: during AP; %: percentage; n: number; SD: standard deviation; p: P-value; RAP: recurrent acute pancreatitis; IQR: interquartile range; CP: chronic pancreatitis; DM: diabetes mellitus in the personal medical history; Hyperlip: hyperlipidemia in the personal medical history.

BAP – HbA1c								р
Groups, HbA1c%	≤6.50		6.51-7.00	7.01-8.00		8.01-9.00	≥9.01	_
n (%)	633 (84.0)		28 (3.7)	32 (4.2)		22 (2.9)	39 (5.2)	
Age (years), mean (SD)	56.50 (16.76)		63.54 (12.20)	58.38 (12.76)		55.23 (11.59)	53.08 (13.86)	0.068
Female, n (%)	260 (41)		13 (46)	10 (31)		4 (18)	11 (28)	0.078
RAP, n (%)	142 (22)		2 (7)	7 (22)		7 (32)	10 (26)	0.561
Episodes, n, median	138; 1.00 (1.00, 2	2.00)	2; 1.00 (1.00, 1.00)	6; 1.00 (1.00, 1.00)		7; 1.00 (1.00, 3.50)	10; 1.50 (1.00, 2.75)	0.845
CP n (%)	48 (8)		3 (11)	3 (9)		2 (9)	2(5)	0 882
DM. n (%)	74 (11.7)		17 (60.7)	28 (87.5)		18 (81.8)	35 (89.7)	< 0.001
Hyperlip, n (%)	56 (9)		4 (16)	6(21)		5 (26)	16 (43)	< 0.001
OAP – on-admission serum glucose							()	D
Groups, mmol/L	<3.99	4-5.99	6.00-7.79	7.80-11.09	11.10-14.99	15.00-19.99	>20.00	r
n (%)	21 (1.0)	468 (22.0)	697 (32.7)	613 (28.8)	221 (10.4)	68 (3.2)	41 (1.9)	
Age (years), mean (SD)	45.52 (19.88)	51.40 (17.23)	56.12 (17.62)	59.92 (15.56)	61.20 (14.81)	58.62 (13.91)	53.05 (15.23)	<0.001
Female, n (%)	12 (57)	208 (44)	294 (42)	266 (43)	96 (43)	23 (34)	14 (34)	0.423
RAP, n (%)	2 (10)	109 (23)	161 (23)	134 (22)	40 (18)	16 (24)	6 (15)	0.216
Episodes, n, median	2; 1.50 (1.25,	102; 1.00 (1.00,	152; 1.00 (1.00,	124; 1.00 (1.00,	35; 1.00 (1.00,	16; 1.00 (1.00,	6; 2.50 (2.00,	0.227
(IQR)	1.75)	2.00)	2.00)	2.00)	2.00)	2.50)	3.75)	
CP, n (%)	0(0)	30 (6)	46 (7)	30 (5)	6 (3)	3 (4)	4 (10)	0.259
DM, n (%)	2 (9.5)	24 (5.1)	62 (8.9)	121 (19.7)	105 (47.5)	49 (72.1)	30 (73.2)	<0.001
Hyperlip, n (%)	2 (13)	41 (10)	67 (11)	100 (19)	40 (21)	15 (25)	18 (50)	<0.001
DAP – peak serum glucose								р
Groups, mmol/L	≤3.99	4.00-5.99	6.00-7.79	7.80-11.09	11.10-14.99	15.00-19.99	$\geq 20.00$	
n (%)	6 (1.0)	187 (31.3)	143 (24.0)	123 (20.6)	79 (13.2)	43 (7.2)	16 (2.7)	
Age (years), mean (SD)	60.50 (19.21)	58.03 (18.97)	57.83 (16.13)	59.54 (15.19)	60.82 (13.24)	57.84 (14.44)	61.50 (14.02)	0.844
Female, n (%)	4 (67)	88 (47)	63 (44)	50 (41)	29 (37)	19 (44)	8 (50)	0.611
RAP, n (%)	3 (50)	37 (20)	34 (24)	27 (22)	19 (24)	8 (19)	3 (19)	0.868
Episodes, n, median	3; 2.00 (1.50,	35; 2.00 (1.00,	30; 1.00 (1.00,	22; 1.00 (1.00,	16; 1.00 (1.00, 3.00	) 7; 1.00 (1.00,	3; 1.00 (1.00,	0.577
(IQR)	2.00)	2.00)	2.00)	3.00)		1.50)	1.50)	
CP, n (%)	0(0)	11 (6)	6 (4)	3 (2)	6 (8)	4 (9)	0 (0)	0.770
DM, n (%)	0(0)	13 (7.0)	17 (11.9)	28 (22.8)	46 (58.2)	29 (67.4)	10 (62.5)	<0.001
Hyperlip, n (%)	2 (33)	18 (12)	21 (18)	19 (17)	23 (31)	13 (33)	6 (40)	<0.001



**Fig. 1.** Etiology distribution. The top row shows the classification of groups. The middle row shows how etiologies within each group (etiologies add up to 100%). The bottom row shows group distribution within each etiology (groups add up to 100%). AP: acute pancreatitis; BAP: before AP; OAP: on-admission with AP; DAP: during AP; HbA1c: hemoglobin A1c; Glu: serum glucose; HTG: hypertriglyceridemia; bil: biliary, alc: alcoholic; comb: combined; idiop: idiopathic.



Fig. 2. AP severity, mortality, length of AP-associated hospital stay, maximal CRP. AP: acute pancreatitis; BAP: before AP; OAP: on-admission with AP; DAP: during AP; HbA1c: hemoglobin A1c; Glu: serum glucose; LOH: length of AP associated hospitalization; max. CRP: maximal C-reactive protein. Some statisctical outliers were removed (not omitted) in case of the LOH and max. CRP boxplots to facilitate interpretation, original figures can be found in the supplementary material (Supplementary Fig. 3).

Group 2 (7.1%) (Fig. 2).

No trends were identified regarding systemic complications (respiratory, renal or heart failure) (p = 0.959); the highest rate of organ failures was found in Group 3 (12.5%) (Fig. 3). On the other hand, an increasing proportion of local complications (peripancreatic fluid, pancreas pseudocyst, pancreas necrosis) were seen with increasing HbA1c (Fig. 4), without statistical significance (p = 0.122).

The binary logistic regression did not show HbA1c to be an independent predictor of mortality (OR = 1.211 (95% CI: 0.859–1.646), p = 0.241) or severity (OR = 1.028 (95% CI: 0.768–1.332), p = 0.843) (Supplementary Tables 8–9). The ROC analysis (Supplementary Fig. 4) showed that HbA1c fails to predict mortality (AUC = 0.545) and poorly predicts the severity of AP (AUC = 0.601).

OAP and DAP: on-admission and peak glucose levels demonstrate a dose-dependent association with worse AP outcomes (severity, mortality, complications, LOH, maximal CRP)

A dose-dependent association was seen between on-admission glucose levels, peak in-hospital glucose levels and: severity (p < 0.001 in both OAP and DAP), mortality (p < 0.001 OAP and DAP), LOH (p < 0.001 OAP and DAP), maximal CRP (p < 0.001 OAP and DAP) (Fig. 2), systemic complications (p < 0.001 OAP and DAP; Fig. 3) and local complications (p < 0.001 OAP and DAP; Fig. 4).



**Fig. 3.** Systemic complications as defined by the revised Atlanta classification and Marshall criteria. The top row shows the classification of groups. The second row shows the percentage of AP cases with organ failure (one or more of lungs, heart and kidney). The third, fourth, and bottom rows show the percentage of AP cases with respiratory, heart and renal failure, respectively, in each group. AP: acute pancreatitis; BAP: before AP; OAP: on-admission with AP; DAP: during AP; HbA1c: hemoglobin A1c; Glu: serum glucose.

The group with a peak in-hospital glucose  $\geq 20 \text{ mmol/L}$  (Group 7) noted the highest severity (37.5% moderate, 25.0% severe), mortality (12.5%), systemic (43.8%) and local complications (62.5%). While similarly, on-admission glucose Group 7 saw the highest rate of local complications (36.6%), Group 6 (serum glucose 15–19.99 mmol/L) had the highest severity (38.2% moderate, 8.8% severe), mortality (8.8%), and rate of systemic complications (14.7%).

The binary logistic regression established both on-admission and peak in-hospital serum glucose to be independently associated with mortality (OR = 1.133 (95% CI: 1.064–1.204), p < 0.001 and OR = 1.089 (95% CI: 1.020–1.161), p = 0.006, respectively) and severity (OR = 1.131 (95% CI: 1.078–1.186), p < 0.001 and OR = 1.093 (95% CI: 1.039–1.152), p < 0.001, respectively) (Supplementary Tables 8–9). The ROC analysis (Supplementary Fig. 4) showed on-admission glucose to be a poor predictor of mortality (AUC = 0.636 for an estimated cut-off of 10.635 mmol/l) and severity (AUC = 0.671 for an estimated cut-off of 9.435 mmol/l). An on-admission glucose >10 mmol/l had an OR of 3.140 (95% CI: 2.106–4.682) for severe AP and an OR of 2.666



**Fig. 4.** Local pancreatic complications as defined by the revised Atlanta classification. The top row shows the classification of groups. The second row shows the percentage of AP cases with local complications (one or more of acute peripancreatic fluid collection, pancreas pseudocyst, pancreas necrosis – no distinction between acute necrotic collection and walled-off necrosis). The third, fourth, and bottom rows show the percentage of AP cases with acute peripancreatic fluid collection, pancreas necrosis, respectively, in each group. AP: acute pancreatitis; BAP: before AP; OAP: on-admission with AP; DAP: during AP; HbA1c: hemoglobin A1c; Glu: serum glucose.

(95% CI: 1.587–4.478) for mortality. The ROC analysis indicated that peak in-hospital glucose is a fair predictor of mortality (AUC = 0.703 for an estimated cut-off of 6.665 mmol/l) and severity (AUC = 0.732 for an estimated cut-off of 7.355 mmol/l). A peak in-hospital glucose >7 mmol/l had an OR of 14.490 (95% CI: 4.443–47.264) for severe AP and an OR of 4.750 (95% CI: 1.370–16.476) for mortality.

Hypoglycemia (Group 1) was associated with a higher rate of organ failure and maximal CRP in both the OAP and DAP subcohorts

and higher severity only in the DAP subcohort. Increased mortality, LOH or local complications were not seen.

## Additional trends, intergroup differences

The results of additional trend tests not discussed in the article's main body and P-values corresponding to intergroup differences for all examined outcomes can be found in the supplementary material (Supplementary Tables 5–7).

## Discussion

Pancreatic inflammation together with cell death are the end steps of an intricate interplay of events - as described by the model of "multiple hits on multiple targets", various signaling pathways are activated resulting in the clinical entity of AP [24]. The main alterations observed inside the acinar cells are the increase in calcium concentration, increase in the number of lysosomes and zymogen granules, leading to mitochondrial dysfunction, endoplasmic reticulum stress and premature trypsinogen activation [3]. These changes can be initiated either by ductal obstruction [25,26] or by the direct acinar effect of various pancreatotoxic agents [27,28]. The most common culprits responsible for pancreatic toxicity are bile acids, alcohol and its metabolites, but fatty acids and fatty acid ethyl esters are also described to initiate the abovementioned pathways in a dose-dependent manner [27,29–32]. And observations go beyond basic science - cohort studies also describe hypertriglyceridemia to be dose-dependently associated with increasing severity and rate of complications [8]. Some drugs as well and smoking are presumed to directly damage the pancreas in a dose dependent manner [33,34].

There is a layered relationship between DM and AP. In severe AP cases with substantial pancreatic necrosis,  $\beta$ -cell loss can lead to the development of DM [35]. But even in moderate and mild AP, impaired β-cell function and insulin resistance are observed in more than 30% of the cases [36], leading to a two-fold risk for developing diabetes after experiencing a single episode [37,38]. At the same time, cohort studies described a 1.5–3 times higher risk of AP among type 2 diabetic individuals – a possible reason behind this are the overlapping risk factors (e.g., obesity and hypertriglyceridemia) [7,39-41]. Another key point in the many intersections between these two conditions is the higher severity of AP in individuals with pre-existing DM. A meta-analysis of cohort studies described a significantly higher risk of complications, intensive care unit (ICU) admission, and mortality as compared to non-DM individuals [42]. In a pancreatitis rat model, streptozotocin-induced hyperglycemia significantly reduced the pancreatic amylase content, which the investigators believed to be due to the direct harmful effects on acinar cells [43]. This indicates that glucose itself could be involved in AP as a potential pancreatotoxic agent. So far, clinical investigations were limited to discussing the role of DM – and neglected the glycemic state.

Our study is the first to focus on how alterations of the glucose homeostasis affect clinically relevant outcomes of AP. We found onadmission and peak in-hospital serum glucose concentrations to have a statistically significant dose-dependent relationship with AP severity, mortality, LOH, maximal CRP, systemic and local complications. Both these variables are independently (accounting for DM, age, gender and etiology) associated with AP severity and mortality, a peak in-hospital glucose >7 mmol/l being associated with a 15 times higher likelihood of severe AP and five times higher likelihood of death. While statistically significant dose-dependency was only identified with LOH and maximal CRP in case of HbA1c, a trend of increasing severity and rate of local complications was also noted. In light of the currently available scientific literature on the matter, these findings strongly suggest that glucose has a direct pancreatotoxic effect.

The main step in glucose-mediated cytotoxicity is the intracellular increase in reactive oxygen species (ROS) [44]. ROS also have a central role in the process of pancreatic inflammation, promoting pathways towards mitochondrial dysfunction, cell death and inflammation in a self-amplifying manner [45,46]. Another way in which glucose could potentially harm the pancreas is via influencing its vasculature. Increased ROS production inside endothelial cells in hyperglycemia causes microvascular endothelial dysfunction by decreasing nitric oxide availability and increasing permeability, leukocyte adhesion and procoagulant activity [47]. Such microvascular disturbances contribute to the inflammation of the ischemia-sensitive pancreas [48].

While we demonstrated dose-dependency and independent association with AP severity and mortality, the whole extent of worsening AP outcomes should not exclusively be attributed to glucose. As hyperglycemia often presents in the context of DM and metabolic syndrome, the prevalence of these conditions accumulates with increasing glucose concentration. Here, an increased rate of hypertriglyceridemia, cholelithiasis and possibly β-cell hypertrophy facilitate the formation of AP [49-52], with hypertriglyceridemia also being associated with a more severe disease course in most studies, although controversy still exists [53–55]. Accordingly, in our cohort, we observed a growing proportion of hypertriglyceridemia with increasing serum glucose and HbA1c values. Severity and mortality however, only increased parallelly with on-admission and peak glucose, but not HbA1c. This might partly be due to the low participant and event numbers in the higher HbA1c groups - 83.9% of cases had HbA1c values below 6.5% - and partly because HbA1c indicates the preceding three months' glucose homeostasis, not necessarily reflecting glucose levels at the time of the AP. All in all, the fact that the proportion of hypertriglyceridemia increased with all three variables, but severity and mortality only did so with serum glucose further reinforces the results of our binary logistic regression, underlining the potential pancreatotoxic nature of hyperglycemia.

Apart from signaling a transient dysregulation, on-admission and especially peak in-hospital hyperglycemia could also be caused by new-onset pancreatogenic DM. A common concern is that AP episodes that are more severe might have a higher likelihood of substantial  $\beta$ -cell death; this could increase the proportion of severe cases among those with high glucose values. However, clinical studies do not support the association between AP severity and newly diagnosed DM [56]. DM and thus pathological serum glucose values can be overrepresented in CP cases, 25–80% of CP patients develop DM [57]. Acute exacerbations in people with CP are known to be less severe [58], possibly decreasing severity and mortality in higher HbA1c and glucose groups, but we found a balanced distribution of CP among groups.

## Strengths and limitations

**Strengths**: To our knowledge, this is the first clinical study focusing on the role of glucose homeostasis in AP, observing the presence of dose-dependency with its clinically important outcomes. We succeeded in demonstrating dose-dependency and independent association with AP severity and mortality for both on-admission and peak in-hospital serum glucose. The data source is a prospective cohort, boasting an impressive number of patients from multiple centers worldwide, enhancing the applicability of our results. As described in the 'Methods' section we also applied a rigorous quality control system, to ensure the validity of our data.

**Limitations**: A limitation to our study is that glucose values cannot be separated from the underlying DM and metabolic comorbidities, possibly influencing outcomes. Nevertheless, we feel that they should not be separated: although more likely in diabetic patients, hyperglycemia can occur and elicit pancreatotoxicity in any subpopulation. Distinguishing exactly what causes the dysregulation in each patient was beyond the scope of this clinically oriented cohort analysis – an interesting question nonetheless for future studies. Another limitation is that DM is overrepresented in the BAP subcohort (23.0% of HbA1c measurements happened in known diabetic participants). Only 26.5% of patients had at least two glucose measurements during their

hospitalization, predominantly those with abnormal on-admission values (44.7%). As mentioned in the 'Methods' section, capillary glucose values (although representing the minority of cases) were also accepted. In an attempt to form groups based on established cut-offs and equal increments, some groups ended up having relatively low participant numbers, weakening statistical power.

## Implications

**Prevention:** Increased HbA1c was associated with higher severity and a higher rate of local complications. Maintaining a normal glucose homeostasis might reduce the risk of these events.

**Prognosis:** Increased on-admission glucose has a dosedependent association with increasing severity, mortality, LOH and complications of AP.

**Prompt treatment:** High peak glucose is dose-dependently associated with a higher rate of severe cases, mortality, systemic complications and increased LOH. Hyperglycemia does not necessarily present on admission, monitoring serum glucose during the course of AP is crucial. Adequate in-hospital control of hyperglycemia could greatly contribute to the treatment of AP.

### Author contributions

AN and AP drafted the original concept, and conducted the majority of data interpretation, AN wrote the majority of the manuscript. MFI helped prepare the raw data for statistical analysis. participated in patient enrolment, supervised data quality and wrote the 'Methods' section. AG and AV conducted the statistical analyses. FI, ÁV, PS, JC, ZS, TT, MP, EF, HJ, KK, IT, DS, GP, ATI, BE, KM, DP, DI, OF, TN, PK, ZM all provided a substantial number of enrolled participants. SV and MF participated in patient enrolment, organized and prepared raw data for statistical analysis. NF acted as a radiological supervisor, evaluating radiological findings and helping interpretation. AS, PH and AP acted as supervisors of the article writing process. All co-authors (1) made contributions to the concept and design of the study as well as the data analysis and interpretation; (2) aided the drafting of the article especially in regards to intellectual content; (3) read and approved the final version of the manuscript.

## **Declaration of competing interest**

Authors declare no conflict of interest.

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## Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.pan.2021.06.003.

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#### A. Nagy, M.F. Juhász, A. Görbe et al.

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