



Pancreatic Enzyme Testing for Acute Pancreatitis

Policy Number: AHS – G2153 – Pancreatic Enzyme Testing for Acute Pancreatitis	Prior Policy Name and Number, as applicable:
Effective Date: 02/01/2023	

POLICY DESCRIPTION | RELATED POLICIES | INDICATIONS AND/OR LIMITATIONS OF COVERAGE | TABLE OF TERMINOLOGY | SCIENTIFIC BACKGROUND | GUIDELINES AND RECOMMENDATIONS | APPLICABLE STATE AND FEDERAL REGULATIONS | APPLICABLE CPT/HCPCS PROCEDURE CODES | EVIDENCE-BASED SCIENTIFIC REFERENCES | REVISION HISTORY

I. Policy Description

Pancreatitis is an inflammation of pancreatic tissue and can be either acute or chronic. Pancreatic enzymes, including amylase, lipase, and trypsinogen can be used to monitor the relative health of the pancreatic tissue. Damage to the pancreatic tissue, including pancreatitis, can result in elevated pancreatic enzyme concentrations whereas depressed enzyme levels are associated with exocrine pancreatic insufficiency (P. A. Banks et al., 2013; Stevens & Conwell, 2020).

II. Related Policies

Policy Number	Policy Title
	Not applicable

III. Indications and/or Limitations of Coverage

Application of coverage criteria is dependent upon an individual's benefit coverage at the time of the request. Medical Policy Statements do not ensure an authorization or payment of services. Please refer to the plan contract (often referred to as the Evidence of Coverage) for the service(s) referenced in the Medical Policy Statement. If there is a conflict between the Medical Policy Statement and the plan contract (i.e., Evidence of Coverage), then the plan contract (i.e., Evidence of Coverage) will be the controlling document used to make the determination. Specifications pertaining to Medicare and Medicaid can be found in Section VII of this policy document

Application of coverage criteria is dependent upon an individual's benefit coverage at the time of the request. If there is a conflict between this Policy and any relevant, applicable government policy [e.g. National Coverage Determinations (NCDs) for Medicare] for a particular member, then the government policy will be used to make the determination. For the most up-to-date Medicare policies and coverage, please visit their search website <u>https://www.cms.gov/medicare-coverage-database/search.aspx</u> or <u>the manual website.</u>

 Measurement of either serum lipase (preferred) OR amylase concentration for the initial diagnosis of acute pancreatitis MEETS COVERAGE CRITERIA in all patients presenting with signs and symptoms of acute pancreatitis* (please see Note 1).





- 2) Measurement of either serum lipase (preferred) **OR** amylase concentration **DOES NOT MEET COVERAGE CRITERIA** in the following situations:
 - a) As part of an ongoing assessment of therapy for acute pancreatitis; OR
 - b) In determining the prognosis of pancreatitis; OR
 - c) In determining the severity or progression of pancreatitis; OR
 - d) More than once per visit; OR
 - e) For the diagnosis, prognosis, or severity of chronic pancreatitis; OR
 - f) As part of ongoing assessment or therapy of chronic pancreatitis
 - g) In asymptomatic nonpregnant individuals during general exam without abnormal findings
- 3) Measurement of the following biomarkers for the diagnosis or assessment of acute pancreatitis, prognosis, and/or determination of severity of acute pancreatitis DOES NOT MEET COVERAGE CRITERIA:
 - a) measurement of both amylase AND serum lipase; OR
 - b) serum or urine trypsin/trypsinogen/TAP (trypsinogen activation peptide)

The following does not meet coverage criteria due to a lack of available published scientific literature confirming that the test(s) is/are required and beneficial for the diagnosis and treatment of a patient's illness.

- 4) Measurement of the following biomarkers for the diagnosis or assessment of acute pancreatitis, prognosis, and/or determination of severity of acute pancreatitis DOES NOT MEET COVERAGE CRITERIA:
 - a) C-Reactive Protein (CRP)
 - b) Interleukin-6 (IL-6)
 - c) Interleukin-8 (IL-8)
 - d) Procalcitonin
- 5) Measurement of urinary amylase concentration for the initial diagnosis of acute pancreatitis **DOES NOT MEET COVERAGE CRITERIA** in all patients presenting with signs and symptoms of acute pancreatitis* (please see Note 1)

*Note 1: Acute Pancreatitis Signs and Symptoms (Vege, 2021a):

- Persistent, severe epigastric pain (that may be in the right upper quadrant for some patients)
- Nausea
- Vomiting
- "Approximately 5 to 10 percent of patients with acute severe pancreatitis may have painless disease and have unexplained hypotension."
- Tender to palpitation of epigastrium
- Abdominal distention
- Hypoactive bowel sounds





- Fever
- Rapid pulse
- Tachypnea
- Hypoxemia
- Hypotension

IV. Table of Terminology

Term	Definition
AACC	American Association for Clinical Chemistry
ABIM	American Board of Internal Medicine
ACCR	Amylase-to-creatinine clearance ratio
ACG	American College of Gastroenterology
AED	Academy For Eating Disorders
AGA	American Gastroenterological Association
AP	Acute pancreatitis
APA	American Pancreatic Association
APA	American Psychiatric Association
APACHE-II	Acute Physiology and Chronic Health Evaluation
ASCP	American Society for Clinical Pathology
AUCs	Area under the curve
BISAP	Bedside index for severity in acute pancreatitis
BUN	Blood urea nitrogen
cCRP	Cardiac CRP
CECT	Contrast-enhanced computed tomography
CLIA '88	Clinical Laboratory Improvement Amendments Of 1988
CMS	Centers For Medicare and Medicaid
СР	Chronic pancreatitis
CPEC	Clinical Practice and Economics Committee
CPT	Current procedural terminology
CRP	C-reactive protein
СТ	Computed axial tomography
CTSI	Computed axial tomography severity index
ED	Eating disorder
ELISA	Enzyme-linked immunoassay
ERCP	Endoscopic retrograde cholangiopancreatography
EUS	Endoscopic ultrasonography
FDA	Food and Drug Administration
GRADE	Grading of recommendations assessment, development, and evaluation
HCPCS	Healthcare common procedure coding system
HMGB1	High Mobility Group Box 1
hsCRP	High sensitivity C-reactive protein
HSROC	Hierarchical summary receiver operating characteristics curve
IAP	International Association of Pancreatology
IL-6	Interleukin-6
IL-8	Interleukin-8





LDH	Lactate dehydrogenase	
LDTs	Laboratory-developed tests	
MODS	Multiorgan dysfunction syndrome	
MRCP	Magnetic resonance cholangiopancreatography	
MRI	Magnetic resonance imaging	
NASPGHAN	North American Society for Pediatric Gastroenterology, Hepatology and Nutrition	
NASPORAN	Pancreas Committee	
PBMCs	Peripheral blood mononuclear cells	
PCT	Procalcitonin	
POC	Point of care	
RIA	Radioimmunoassay	
SIRS	Systemic inflammatory response syndrome	
s-isoform	Salivary glands	
SPINK	Serine protease inhibitor	
ТАР	Trypsinogen activation peptide	
ULN	Upper limit of normal	
URL	Upper limit of reference interval	
WSES	The World Society of Emergency Surgery	

V. Scientific Background

Acute Pancreatitis

Acute pancreatitis (AP) is inflammation of the pancreatic tissue that can range considerably in clinical manifestations. Due to the lack of consensus in diagnosing, characterizing, and treating AP, an international group of researchers and practitioners convened in Atlanta in 1992 to write a clinically based classification system for AP, which is now commonly referred to as the Atlanta convention or Atlanta classification system (Bradley & Iii, 1993). The Atlanta classification system was then revised in 2012 (P. A. Banks et al., 2013). For the diagnosis of AP, two of the three following criteria must be present: "(1) abdominal pain consistent with acute pancreatitis (acute onset of a persistent, severe, epigastric pain often radiating to the back); (2) serum lipase activity (or amylase activity) at least three times greater than the upper limit of normal; and (3) characteristic findings of acute pancreatitis on contrast-enhanced computed tomography (CECT) and less commonly magnetic resonance imaging (MRI) or transabdominal ultrasonography" (italics emphasized by the manuscript's authors) (P. A. Banks et al., 2013). This two-of-three criterion is recommended for diagnostic use by several professional societies (P. Banks & Freeman, 2006; Guidelines, 2013; S. Tenner, Baillie, DeWitt, Vege, & American College of, 2013). AP can be characterized by two temporal phases, early or late, with degrees of severity ranging from mild (with no organ failure) to moderate (organ failure less than 48 hours) to severe (where persistent organ failure has occurred for more than 48 hours). The two subclasses of AP are edematous AP and necrotizing AP. Edematous AP is due to inflammatory edema with relative homogeneity whereas necrotizing AP displays necrosis of pancreatic and/or peripancreatic tissues (P. A. Banks et al., 2013). The figure below from Bollen, Hazewinkel, and Smithuis (2015) outlines the revised Atlanta classification system of AP:





	Acute Pancreatitis				
	Two phases	Early	Late		
		1st week	After 1st week		
2	Severity	Mild	Moderate Severe		Severe
		No organ failure	Organ failur less than 48		Organ failure longer than 48 h
	Two types	Oedematous Necro		Necrotizing	
Co	omplications	< 4 wk : acute peripand > 4 wk: pseudocyst	reatic collection		rk : acute necrotic collection rk : walled-off necrosis

Chronic Pancreatitis

Chronic pancreatitis (CP) is also an inflammation of the pancreatic tissue. The two hallmarks of CP are severe abdominal pain and pancreatic insufficiency (Freedman, 2020). Alcohol-induced chronic pancreatitis (or alcohol pancreatitis) accounts for 60-70% of all cases of CP. CP can be classified in one of nine types (Wilson & Smith, 2015):

- Alcoholic
- Autoimmune (idiopathic)
- Due to trauma
- Inherited factors
- Congenital
- Due to hyperparathyroidism
- Hyperlipidemic
- Due to cystic fibrosis
- Due to protein-energy malnutrition

The endocrine system is comprised of several glands which secrete hormones directly into the bloodstream to regulate many different bodily functions. On the other hand, the exocrine system is comprised of glands which secrete products through ducts, rather than directly into the bloodstream. CP affects both the endocrine and exocrine functions of the pancreas. Fibrogenesis occurs within the pancreatic tissue due to activation of pancreatic stellate cells by toxins (for example, those from chronic alcohol consumption) or cytokines from necroinflammation. Measuring the serum levels of amylase, lipase, and/or trypsinogen is not helpful in diagnosing CP since not every CP patient experiences acute episodes, the relative serum concentrations may be either decreased or unaffected, and the sensitivities of the tests are not enough to distinguish reduced enzyme levels (Witt, Apte, Keim, & Wilson, 2007). The best method to diagnose CP is to histologically analyze a pancreatic biopsy, but this invasive procedure is not always the most practical so "the next best diagnostic methods to demonstrate changes consistent with chronic pancreatitis are computed axial tomography (CT), magnetic resonance





cholangiopancreatography (MRCP), endoscopic retrograde cholangiopancreatography (ERCP) and endoscopic ultrasonography (EUS)" (Wilson & Smith, 2015). Previously, ERCP was commonly used to diagnose CP, but the procedure can cause post-ERCP pancreatitis. Genetic factors are also implicated in CP, especially those related to trypsin activity, the serine protease inhibitor SPINK 1, and cystic fibrosis (Borowitz, Grant, & Durie, 1995; Wilson & Smith, 2015; Witt et al., 2007).

Amylase

Amylase is an enzyme produced predominantly in the salivary glands (s-isoform) and the pancreas (pisoform or p-isoamylase) and is responsible for the digestion of polysaccharides, cleaving at the internal $1 \rightarrow 4$ alpha linkage. Up to 60% of the total serum amylase can be of the s-isoform. The concentration of total serum amylase as well as the pancreatic isoenzyme increase following pancreatic injury or inflammation (Basnayake & Ratnam, 2015; Vege, 2020). Even though the serum concentration of the pancreatic diagnostic enzymes, including amylase, lipase, elastase, and immunoreactive trypsin all increase within 24 hours of onset of symptomology, amylase is the first pancreatic enzyme to return to normal levels so the timing of testing is of considerable importance for diagnostic value (Basnayake & Ratnam, 2015; Ventrucci et al., 1987; Yadav, Agarwal, & Pitchumoni, 2002). The half-life of amylase is 12 hours since it is excreted by the kidneys, so its clinical value decreases considerably after initial onset of AP. The etiology of the condition can also affect the relative serum amylase concentration. In up to 50% of AP instances due to hypertriglyceridemia (high blood levels of triglycerides), the serum amylase concentration falls into the normal range, and normal concentrations of amylase has been reported in cases of alcohol-induced AP (Basnayake & Ratnam, 2015; Quinlan, 2014); in fact, one study shows that 58% of the cases of normoamylasemic AP was associated with alcohol use (Clavien et al., 1989). Elevated serum amylase concentrations also can occur in conditions other than AP, including hyperamylasemia (excess amylase in the blood) due to drug exposure (Ceylan, Evrensel, & Önen Ünsalver, 2016; S. Liu et al., 2016), bulimia nervosa (Wolfe, Jimerson, Smith, & Keel, 2011), leptospirosis (Herrmann-Storck et al., 2010), and macroamylasemia (Vege, 2020). Serum amylase levels are often significantly elevated in individuals with bulimia nervosa due to recurrent binge eating episodes (Wolfe et al., 2011).

Macroamylasemia is a condition where the amylase concentration increases due to the formation of macroamylases, complexes of amylase with immunoglobulins and/or polysaccharides. Macroamylasemia is associated with other disease pathologies, "including celiac disease, HIV infection, lymphoma, ulcerative colitis, rheumatoid arthritis, and monoclonal gammopathy." Suspected macroamylasemia in instances of isolated amylase elevation can be confirmed by measuring the amylase-to-creatinine clearance ratio (ACCR) since macroamylase complexes are too large to be adequately filtered. Normal values range from 3-4% with values of less than 1% supporting the diagnosis of macroamylasemia. ACCR itself is not a good indicator of AP since low ACCR is also exhibited in diabetic ketoacidosis and severe burns (Vege, 2020). Hyperamylasemia is also seen in other extrapancreatic conditions, such as appendicitis, salivary disease, gynecologic disease, extra-pancreatic tumors, and gastrointestinal disease (Terui et al., 2013; Vege, 2020). Gullo's Syndrome (or benign pancreatic hyperenzymemia) is a rare condition that also exhibits high serum concentrations of pancreatic enzymes without showing other signs of pancreatitis (Kumar, Ghosh, Tandon, & Sahoo, 2016). No correlation has been found between the concentration of serum amylase and the severity or prognosis of AP (Lippi, Valentino, & Cervellin, 2012).

Urinary amylase and peritoneal amylase concentrations can also be measured. Rompianesi et al. (2017) reviewed the use of urinary amylase and trypsinogen as compared to serum amylase and serum lipase testing. The authors note that "with regard to urinary amylase, there is no clear-cut level beyond which





someone with abdominal pain is considered to have acute pancreatitis." Three studies regarding urinary amylase were reviewed —each with 134-218 participants—and used the hierarchical summary receiver operating characteristics curve (HSROC) analysis to compare the accuracy of the studies. Results showed that "the models did not converge" and the authors concluded that "we were therefore unable to formally compare the diagnostic performance of the different tests" (Rompianesi et al., 2017).

Another study investigated the use of peritoneal amylase concentrations for diagnostic measures and discovered that patients with intra-abdominal peritonitis had a mean peritoneal amylase concentration of 816 U/L (142-1746 U/L range), patients with pancreatitis had a mean concentration of 550 U/L (100-1140 U/L range), and patients with other "typical infectious peritonitis" had a mean concentration of 11.1 U/L (0-90 U/L range). Conclusions state "that peritoneal fluid amylase levels were helpful in the differential diagnosis of peritonitis in these patients" and that levels >100 U/L "differentiated those patients with other intra-abdominal causes of peritonitis from those with typical infectious peritonitis" (Burkart, Haigler, Caruana, & Hylander, 1991). The authors do not state if intraperitoneal amylase is specifically useful in diagnosing AP.

Lie et al. (2021) conducted a retrospective cohort study to evaluate whether serum amylase and lipase could serve as a biomarker to predict pancreatic injury in 79 critically ill children who died of different causes. Through autopsy investigation, the subjects were divided into pancreatic injury group and non-pancreatic injury group. 41 patients (51.9%) exhibited pathological changes of pancreatic injury. Levels of lactate, erythrocyte sedimentation rate, alanine transaminase, aspartate transaminase, and troponin-I in the pancreatic injury group were significantly higher than that in the noninjury group. "Multivariable logistic regression analysis showed that serum amylase, serum lipase, and septic shock were significantly associated with the occurrence rate of pancreatic injury." Therefore, the authors conclude that "serum amylase and lipase could serve as independent biomarkers to predict pancreatic injury in critically ill children" (P. Liu et al., 2021).

In a prospective case control study, Judal (2022) investigated urinary amylase levels for diagnosis of acute pancreatitis. One major challenge with measurement of serum amylase is its short half-life which returns to normal levels within a short period of time. This study enrolled 100 patients (50 healthy and 50 with acute pancreatitis) who were measured for serum amylase, serum lipase, and urinary amylase. There was a statistically significant increase in the serum amylase, lipase, and urinary amylase mean values of patients with AP. "Serum amylase had the highest sensitivity (100%) and serum lipase had the highest specificity (96.53%). The sensitivity and specificity of urinary amylase was found to be 97.25% and 91.47% respectively" (Judal, 2022). The authors conclude that urinary amylase is a convenient and sensitive test for diagnosis.

Lipase (Pancreatic Lipase or Pancreatic Triacylglycerol Lipase)

Pancreatic lipase or triacylglycerol lipase (herein referred to as "lipase") is an enzyme responsible for hydrolyzing triglycerides to aid in the digestion of fats. Like amylase, lipase concentration increases shortly after pancreatic injury (within 3-6 hours). However, contrary to amylase, serum lipase concentrations remain elevated for 1-2 weeks after initial onset of AP since lipase can be reabsorbed by the kidney tubules (Lippi et al., 2012). Moreover, the pancreatic lipase concentration is 100-fold higher than the concentration of other forms of lipases found in other tissues such as the duodenum and stomach (Basnayake & Ratnam, 2015). Both the sensitivity and the specificity of lipase in laboratory testing of AP are higher than that of amylase (Yadav et al., 2002). A study by Coffey, Nightingale, and Ooi (2013) found "an odds ratio of 7.1 (95% confidence interval 2.5-20.5; P<0.001) for developing severe AP" in patients ages 18 or younger when the serum lipase concentration is at least 7-fold higher than





upper limit of normal. However, in general, elevated serum lipase concentration is not used to determine the severity or prognosis of AP (Ismail & Bhayana, 2017). Hyperlipasemia can also occur in other conditions such as Gullo's Syndrome (Kumar et al., 2016). The use of lipase to determine etiology of AP is of debate. A study by Levy et al. (2005) reports that lipase alone cannot be used to determine biliary cause of AP whereas other studies have indicated that a ratio of lipase-to-amylase concentrations ranging from 2:1 to more than 5:1 can be indicative of alcohol-induced AP (Gumaste, Dave, Weissman, & Messer, 1991; Ismail & Bhayana, 2017; Pacheco & Oliveira, 2007; S. M. Tenner & Steinberg, 1992).

The review by Ismail and Bhayana (2017) included a summary table (Table 1 below) comparing various studies concerning the use of amylase and lipase for diagnosis of AP as well as a table (Table 2 below) comparing the cost implication of the elimination of double-testing for AP.

Table 1: Summary of numerous studies comparing lipase against amylase (URL – Upper Limit of Reference interval, AP – Acute Pancreatitis).

Design and reference	Participant (patients with	Threshold	Results		Conclusion
	abdominal pain/AP)		Serum lipase	Serum amylase	
Prospective study [56]	384/60	Two times URL	Diagnostic accura are > 95% for bot	• •	No difference between amylase and lipase in diagnosing AP
Prospective study [57]	306/48	Serum lipase > 208 U/L Serum amylase > 110 U/L	92% sensitivity 87% specificity 94% Diagnostic accuracy	93% sensitivity 87% specificity 91% Diagnostic accuracy	Both tests are associated with AP, but serum lipase is better than amylase
Prospective study [58]	328/51	Serum lipase: > 208 U/L (Day 1) > 216 U/L (Day 3) Serum amylase: > 176 U/L > 126 U/L (Day 3)	Day 1: 64 % Sensitivity 97% Specificity Day 3: 55% Sensitivity 84% Specificity	Day 1: 45 % Sensitivity 97% Specificity Day 3: 35% Sensitivity 92% Specificity	Serum lipase is better at diagnosing early and late AP
Retrospective study [63]	17,531/320 *49 had elevated lipase only	Serum lipase > 208 U/L Serum amylase > 114 U/L	90.3% Sensitivity 93.6% Specificity	78.7% Sensitivity 92.6% Specificity	Serum lipase is more accurate marker for AP
Cohort study [2]	1,520/44	Three times URL	64% Sensitivity 97% Specificity	50% Sensitivity 99% Specificity	Serum lipase is preferable to use in comparison to





					amylase alone or both tests
Retrospective study [59]	3451/34 *33 patients had elevated amylase and 50 had elevated lipase only	Three or more times URL	95.5% Sensitivity 99.2% Specificity	63.6% Sensitivity 99.4% Specificity	Both enzymes have good accuracy, but lipase is more sensitive than amylase
Cohort study [60]	151/117 *6 patients with gallstone-induced and 5 patients with alcohol- induced AP had elevated lipase only	Three times URL	96.6% Sensitivity 99.4% Specificity	78.6% Sensitivity 99.1% Specificity	Lipase is more sensitive in diagnosing AP and using it alone would present a substantial cost saving on health care system
Prospective study [61]	476/154 *58 patients had a normal amylase level	Three times URL	91% Sensitivity 92% Specificity	62% Sensitivity 93% Specificity	Lipase is more sensitive than amylase and should replace amylase in diagnosis of AP
Cohort study [62]	50/42 *8 patients had elevated lipase only	Three times URL	100% Sensitivity	78.6% Sensitivity	Lipase is a better choice than amylase in diagnosis of AP

This table is a list of individual studies examining the specificity and sensitive of serum lipase and serum amylase in diagnosing AP. In each of the listed studies except one, the authors concluded that serum lipase is better than serum amylase for AP. The only outlier used a lower threshold in considering enzyme elevation; in particular, two times the upper limit of reference interval (URL) was used whereas the Atlanta classification system recommends at least three times the URL to determine enzyme elevation (Ismail & Bhayana, 2017).

Table 2: Summary of studies exploring the cost implication associated with eliminating amylase test.

Design and Reference	Costs	Volume of test	Results
Cohort study (UK) [2]	Amylase costs £1.94 Lipase cost £2.50	1383 request for 62 days costing £6136 for both tests	 Testing lipase only will result in cost saving
Cohort study (UK) [60]	Single amylase or lipase cost about £0.69 each Cost of both measured together were £0.99.	2979 requests costing £2949.21	 Measuring lipase would save health care system an estimate of £893.70 per year





Prospective study (US) [71]	Patients charged \$35 for either lipase or amylase	618 co-ordered both lipase and amylase	Amylase test was removed from common order sets in the electronic medical record
			 Reduced the co-ordering of lipase and amylase to 294
			 Overall saving of \$135,000 per year

This table specifically outlines studies that compared the financial cost of the serum amylase and serum lipase tests for diagnosing AP. All three studies show cost savings if only lipase concentration is used. In fact, one study by researchers in Pennsylvania resulted in the removal of the amylase test "from common order sets in the electronic medical record" (Ismail & Bhayana, 2017).

Furey, Buxbaum, and Chambliss (2020) compared amylase and lipase ordering patterns for patients with AP. A total of 438 individuals were included in this study. The researchers noted that "All patients had at least one lipase ordered during their admission, and only 51 patients (12%) had at least one amylase ordered. On average, lipase was elevated 5 times higher above its respective upper reference limit than amylase at admission" (Furey et al., 2020). Further, patients undergoing a laparoscopic cholecystectomy (gallbladder removal) were more likely to have amylase ordered. These results showed that in 88% of patients with AP, amylase measurement was not necessary; moreover, "Of patients for whom amylase was ordered, it was common for these patients to be those referred to surgical procedures, possibly because amylase normalization may be documented faster than that of lipase" (Furey et al., 2020).

In a retrospective cross-sectional study by El Halabi et al. (2019), the clinical utility and economic burden of routine serum lipase examination in the emergency department was observed. From 24,133 adult patients admitted within a 12-month period, serum lipase levels were ordered for 4,976 (20.6%) patients. Of those 614 (12.4%) who had abnormal lipase levels, 130 of those patients were above the diagnostic threshold for acute pancreatitis (>3 times the ULN) and 75 patients had confirmed diagnosis of acute pancreatitis. 1,890 patients had normal no abdominal pain or history of acute pancreatitis, but 251 of these patients were tested for lipase levels, leading to a total cost of \$51,030. These results triggered unneeded cross-sectional abdominal imaging in 61 patients and unwarranted gastroenterology consultation in 3 patients, leading to an additional charge of \$28,975. The authors conclude that "serum lipase is widely overutilised in the emergency setting resulting in unnecessary expenses and investigations" (El Halabi et al., 2019).

Liu et al. (2021) studied the use of serum amylase and lipase for the prediction of pancreatic injury in critically ill children admitted to the PICU. 79 children who died from different cases were studied from autopsy and it was found that 41 of these patients had pathological signs of pancreatic injury. Multivariable logistic regression analysis showed that serum amylase, serum lipase, and septic shock were significantly associated with the occurrence rate of pancreatic injury. Serum amylase was measured with 53.7% sensitivity, 81.6% specificity, cut off value of 97.5, and AUC of 0.731. Serum lipase was measured with 36.6% sensitivity, 92.1% specificity, cut off value of 61.1, and AUC of 0.727. The authors conclude that "serum amylase and lipase could serve as independent biomarkers to predict pancreatic injury in critically ill children" (P. Liu et al., 2021).

Trypsin/Trypsinogen/TAP





Trypsin is a protease produced by the pancreatic acinar cells. It is first synthesized in its zymogen form, trypsinogen, which has its N-terminus cleaved to form the mature trypsin. Pancreatitis can result in blockage of the release of the proteases while their synthesis continues. This increase in both intracellular trypsinogen and cathepsin B, an enzyme that can cleave the trypsinogen activation peptide (TAP) from the zymogen to form mature trypsin, results in a premature intrapancreatic activation of trypsin. This triggers a release of both trypsin and TAP extracellularly into the serum and surrounding peripancreatic tissue. Due to the proteolytic nature of trypsin, this response can result in degradation of both the pancreatic and peripancreatic tissues (i.e., necrotizing AP) (Vege, 2021b; Yadav et al., 2002). Trypsin activity "is critical for the severity of both acute and chronic pancreatitis" (Zhan et al., 2019). When the intracellular activity of trypsin escalates, an increase is also reflected in the number of pancreatitis cases overall, as well as in the severity of these cases (Sendler & Lerch, 2020).

Since trypsinogen is readily excreted, a urine trypsinogen-2 dipstick test has been developed (Actim Pancreatitis test strip from Medix Biochemica), which has a reported specificity of 85% for severe AP within 24 hours of hospital admission (Lempinen et al., 2001). Another study reported that the trypsinogen-2 dipstick test has a specificity of 95% and a sensitivity of 94% for AP, which is higher than a comparable urine test for amylase (Kemppainen et al., 1997). As of 2020, the FDA has not approved the use of the trypsinogen-2 dipstick test for the detection or diagnosis of AP. Clinical trials are underway in the United States (Eastler, 2017). The use of TAP for either a diagnostic or prognostic tool is of debate (Lippi et al., 2012).

The study by Neoptolemos et al. (2000) reported that a urinary TAP assay had a 73% specificity for AP. However, another study using a serum TAP methodology reported a 23.5% sensitivity and 91.7% specificity for AP and concluded that "TAP is of limited value in assessing the diagnosis and the severity of acute pancreatic damage" (Pezzilli et al., 2004).

Yasuda et al. (2019) completed a multicenter study in Japan which measured the usefulness of the rapid urinary trypsinogen-2 dipstick test and levels of urinary trypsinogen-2 and TAP concentration as prognostic tools for AP. A total of 94 patients participated in this study from 17 medical institutions between April 2009 and December 2012. The researchers determined that "The trypsinogen-2 dipstick test was positive in 57 of 78 patients with acute pancreatitis (sensitivity, 73.1%) and in 6 of 16 patients with abdominal pain but without any evidence of acute pancreatitis (specificity, 62.5%)" (Yasuda et al., 2019). Further, both TAP and urinary trypsinogen-2 levels were significantly higher in patients with extrapancreatic inflammation. The authors concluded that the urinary trypsinogen-2 dipstick test is a useful tool for AP diagnoses.

Simha at al. (2021) studied the utility of POC urine trypsinogen dipstick test for diagnosing AP in an emergency unit. Urine trypsinogen dipstick test (UTDT) was performed in 187 patients in which 90 patients had AP. UTDT was positive in 61 (67.7%) of the 90 AP patients. In the 97 non pancreatitis cases, UTDT was positive in 9 of those cases (9.3%). The sensitivity and specificity of UTDT for acute pancreatitis was 67.8% and 90.7%, respectively. The authors conclude that although it is a great and convenient possibility as a POC test, "the low sensitivity of UTDT could be a concern with its routine use" (Simha et al., 2021).

Other Biochemical Markers (CRP, Procalcitonin, IL-6, IL-8)

AP results in the activation of the immune system. Specific markers including C-reactive protein (CRP), procalcitonin, interleukin-6 (IL-6), and interleukin-8 (IL-8) have been linked to AP (Toouli et al., 2002; Vege, 2021a; Yadav et al., 2002). CRP is a nonspecific marker for inflammation that takes 48-72 hours to





reach maximal concentration after initial onset of AP but is reported to have a specificity of 93% in detecting pancreatic necrosis. CRP can be used in monitoring the severity of AP; however, imaging techniques, including CT, and evaluative tools, such as the APACHE-II (acute physiology and chronic health evaluation) test, are preferred methods (Guidelines, 2013; Quinlan, 2014).

Procalcitonin is the inactive precursor of the hormone calcitonin. Like CRP, procalcitonin has been linked to inflammatory responses, especially in response to infections and sepsis. Procalcitonin levels are elevated in AP and are significantly elevated (\geq 3.5 ng/mL for at least two consecutive days) in cases of AP associated with multiorgan dysfunction syndrome (MODS) (Rau et al., 2007). Moreover, the elevated procalcitonin levels decrease upon treatment for AP; "however, further research is needed in order to understand how these biomarkers can help to monitor inflammatory responses in AP" (Simsek et al., 2018).

The concentration of inflammatory cytokines IL-6 and IL-8 become elevated in AP with a maximal peak within the first 24 hours after initial onset of AP (Yadav et al., 2002). One study by Jakkampudi et al. (2017) shows that IL-6 and IL-8 are released in a time-dependent manner after injury to the pancreatic acinar cells. This, in turn, activated the peripheral blood mononuclear cells (PBMCs), which propagate acinar cell apoptosis that results in further release of cytokines to increase the likelihood of additional cellular damage.

A study conducted by Khanna et al. (2013) compares the use of biochemical markers, such as CRP, IL-6, and procalcitonin, in predicting the severity of AP and necrosis to that of the clinically used evaluative tools, including the Glasgow score and APACHE-II test. Their results indicate that CRP has a sensitivity and specificity of 86.2% and 100%, respectively, for severe AP and a sensitivity and specificity of 100% and 81.4%, respectively, for pancreatic necrosis. These scores are better than those reported for the clinical evaluative tools (see table below). IL-6 also shows an increase in both sensitivity and specificity; however, the values for procalcitonin are considerably lower than either CRP or IL-6 in all parameters (Khanna et al., 2013).

Data from	<u>Severe AP</u>		<u>Pancreatic r</u>	<u>ecrosis</u>
(Khanna et al., 2013)	Sensitivity	Specificity	Sensitivity	Specificity
Glasgow	71.0	78.0	64.7	63.6
APACHE-II	80.6	82.9	64.7	61.8
CRP	86.2	100	100	81.4
IL-6	93.1	96.8	94.1	72.1
Procalcitonin	86.4	75.0	78.6	53.6

Another study by Hagjer and Kumar (2018) compared the efficacy of the bedside index for severity in acute pancreatitis (BISAP) scoring system to CRP and procalcitonin shows that CRP is not as accurate for prognostication as BISAP. BISAP has AUCs for predicting severe AP and death of 0.875 and 0.740, respectively, as compared to the scores of CRP (0.755 and 0.693, respectively). Procalcitonin, on the other hand, had values of 0.940 and 0.769 for predicting severe AP and death, respectively. The authors concluded that it "is a promising inflammatory marker with prediction rates similar to BISAP" (Hagjer & Kumar, 2018).





Li, Wang, Cai, and Liu (2018) completed a meta-analysis to determine the relationship between high mobility group box 1 (HMGB1), interleukin-6 (IL-6) and AP. HMGB1 protein is a nuclear protein with several different purposes depending on its location (Yang, Wang, Chavan, & Andersson, 2015). These researchers analyzed data from 27 different studies comprised of 1908 of participants (896 with mild AP, 700 with severe AP and 312 healthy controls). Overall, serum HMGB1 and IL-6 levels were higher in patients with both severe and mild AP compared to controls; further, and serum HMGB1 and IL-6 levels were significantly higher in patients with severe AP than mild AP (Li et al., 2018). The authors concluded that serum HMGB1 and IL-6 levels "might be used as effective indicators for pancreatic lesions as well as the degree of inflammatory response" and that both HMGB1 and IL-6 are closely correlated with pancreatitis severity.

Tian et al. (2020) studied the diagnostic value of C-reactive protein (CRP), procalcitonin (PCT), IL-6, and lactate dehydrogenase (LDH) in patients with severe acute pancreatitis. 153 patients were divided into the mild acute pancreatitis group (81) and severe pancreatitis group (72). Significant differences in the values of these enzymes were found between both groups. The sensitivity, specificity, and AUC were determined as seen in the chart below. The AUC of combined detection of CRP, PCT, IL-6 and LDH was 0.989. The authors conclude that "the combined detection of CRP, PCT, IL-6 and LDH has a high diagnostic value for judging the severity of acute pancreatitis" (Tian et al., 2020).

Enzyme	Sensitivity	Specificity	AUC
CRP	55.6%	73%	0.637
РСТ	77.8%	94%	0.929
IL-6	80.2%	85%	0.886
LDH	82.7%	96%	0.919

In a retrospective cohort study, Wei investigated the predictive value of serum cholinesterase (ChE) in the mortality of acute pancreatitis. 692 patients were enrolled in the study and were divided into the ChE-low group (378 patients) or ChE-normal group (314 patients). Mortality was significantly different in two groups (10.3% in ChE-low vs. 0.0% ChE- normal) and organ failure also differed (46.6% ChE-low vs. 8.6% ChE-normal). The area under the curve of serum ChE was 0.875 and 0.803 for mortality and organ failure, respectively. The authors conclude that "lower level of serum ChE was independently associated with the severity and mortality of AP" (Wei et al., 2022).

VI. Guidelines and Recommendations

International Association of Pancreatology (IAP) and the American Pancreatic Association (APA)

In 2012, a joint conference between the International Association of Pancreatology (IAP) and the American Pancreatic Association (APA) convened to address the guidelines for the management of acute pancreatitis. This conference made their recommendations using the Grading of Recommendations Assessment, Development and Evaluation (GRADE) system. The guidelines are detailed with 38 recommendations covering 12 different topics, ranging from diagnosis to predicting severity of disease to timing of treatments. As concerning the diagnosis and etiology of AP, the associations conclude with "GRADE 1B, strong agreement" that the definition of AP follow the Atlanta classification system where at least two of the following three criteria are evident—the clinical manifestation of upper abdominal pain, the laboratory testing of serum amylase or serum lipase where levels are >3 times the upper limit





of normal values, and/or the affirmation of pancreatitis using imaging methods. They specifically did not include the trypsinogen-2 dipstick test in their recommendations "because of its presumed limited availability." One question addressed by the committee was the continuation of oral feeding being withheld for patients until the lab serum tests returned within normal values. With a GRADE 2B, strong agreement finding, they conclude that "it is not necessary to wait until pain or laboratory abnormalities completely resolve before restarting oral feeding." No specific discussion on the preference of either serum amylase or lipase is included within the guidelines as well as no discussion of the use of either serum test beyond initial diagnosis of AP (i.e., no continual testing for disease monitoring is included). Furthermore, no discussion concerning the use of urinary or peritoneal amylase concentrations for AP.

With regards to CRP and/or procalcitonin, the IAP/APA does not address the topic in any detail. As part of their recommendation (GRADE 2B) concerning the best score or marker to predict the severity of AP, they state "that there are many different predictive scoring systems for acute pancreatitis..., including single serum markers (C-reactive protein, hematocrit, procalcitonin, blood urea nitrogen), but none of these are clearly superior or inferior to (persistent) SIRS", which is Systemic inflammatory response syndrome. Moreover, in response to their recommendation for admission to an intensive care unit in AP (GRADE 1C), they state that "the routine use of single markers, such as CRP, hematocrit, BUN or procalcitonin alone to triage patients to an intensive care setting is not recommended."

American Gastroenterological Association (AGA)

The Clinical Practice and Economics Committee (CPEC) of the American Gastroenterological Association (AGA) Institute released the AGA Institute Medical Position Statement on Acute Pancreatitis as approved by the AGA Institute Governing Board in 2007 to address differences in the recommendations of various national and international societies concerning AP. Within their recommendations, they address the necessity of timeliness in the applicability of serum amylase and/or serum lipase testing. Per their recommendations, either serum amylase or serum lipase should be tested within 48 hours of admission. AP is consistent with amylase or lipase levels greater than 3 times the upper limit of the normal value. They specifically state that the "elevation of lipase levels is somewhat more specific and is thus preferred." The AGA guidelines do not address the use of either urinary or peritoneal concentrations of amylase in AP. Also, any patient presenting symptoms of unexplained multiorgan failure or systemic inflammatory response syndrome should be tested for a possible AP diagnosis. Concerning etiology of the phenotype, they suggest that upon admission, "all patients should have serum obtained for measurement of amylase or lipase level, triglyceride level, calcium level, and liver chemistries." Invasive evaluation, such as endoscopic retrograde cholangiopancreatography (ERCP), should be avoided for patients with a single occurrence of AP. The only mention of CRP in their guidelines is in the section concerning the severity (and not the diagnosis of) AP. "Laboratory tests may be used as an adjunct to clinical judgment, multiple factors scoring systems, and CT to guide clinical triage decisions. A serum Creactive protein level >150 mg/L at 48 hours after disease onset is preferred."

In 2018, the AGA published guidelines on the initial management of AP. These guidelines state that "The diagnosis of AP requires at least 2 of the following features: characteristic abdominal pain; biochemical evidence of pancreatitis (ie, amylase or lipase elevated >3 times the upper limit of normal); and/or radiographic evidence of pancreatitis on cross-sectional imaging" (Crockett, Wani, Gardner, Falck-Ytter, & Barkun, 2018).





American College of Gastroenterology (ACG)

The ACG released guidelines concerning AP in both 2006 and 2013. Both sets of guidelines recommend the use of the Atlanta classification system regarding the threshold of either serum amylase or serum lipase levels in the diagnosis of AP (i.e., greater than three times the upper limit of normal range). Both sets of guidelines state that the standard diagnosis is meeting at least two of the three criteria as stated in the revised Atlanta classification system.

The 2006 guidelines discuss the differences between serum amylase and lipase in greater detail. First, although both enzymes can be elevated in AP, the sensitivity and half-life of lipase are more amenable for diagnosis since the levels of lipase remain elevated longer than those of amylase. These guidelines also make note that "it is usually not necessary to measure both serum amylase and lipase" and that "the daily measurement of serum amylase or lipase after the diagnosis of acute pancreatitis has limited value in assessing the clinical progress of the illness." These guidelines discuss the possibility of elevated amylase levels due to causes other than AP, including but not limited to macroamylasemia, whereas the serum levels of lipase are unaffected by these conditions.

The 2013 guidelines do not explicitly state a preference of the serum lipase over serum amylase test in the diagnosis of AP. They also state that lipase levels can be elevated in macrolipasemia as well as certain nonpancreatic conditions, "such as renal disease, appendicitis, cholecystitis, and so on." Neither set of guidelines address the use of either urinary or peritoneal amylase in AP. The 2006 guidelines list other diagnostic tests, including the trypsin/trypsinogen tests as well as serum amyloid A and calcitonin but do not address them further given their limited availability at that time whereas the 2013 guidelines state that, even though other enzymes can be used for diagnostics, "none seems to offer better diagnostic value than those of serum amylase and lipase." They even state that "even the acute-phase reactant C-reactive protein (CRP) the most widely studied inflammatory marker in AP, is not practical as it takes 72h to become accurate."

UK Working Party on Acute Pancreatitis

The UK Working Party on Acute Pancreatitis consists of a consortium of the British Society of Gastroenterology, Association of Surgeons of Great Britain and Ireland, Pancreatic Society of Great Britain and Ireland, and the Association of Upper GI Surgeons of Great Britain and Ireland. The recommendation by the UK Working Party is that "although amylase is widely available and provides acceptable accuracy of diagnosis, where lipase estimation is available it is preferred for the diagnosis of acute pancreatitis (recommendation grade A)." One contrast of the guidelines of the UK Working Party as compared to other professional societies is the relative threshold of the serum concentrations of pancreatic enzymes. Rather than use the >3 times the upper limit of the normal concentrations of either amylase or lipase as stated in the Atlanta classification system, the UK Working Party's guidelines state that AP diagnosis "should not rely on arbitrary limits of values 3 or 4 times greater than normal, but values should be interpreted in light of the time since the onset of abdominal pain." These elevated serum levels as well as the clinical abdominal symptoms are the "cornerstones of diagnosis." They do not address the frequency of serum enzyme testing or the use of trypsin/trypsinogen-based tests, urinary amylase, or peritoneal amylase.





American Board of Internal Medicine (ABIM), American Society for Clinical Pathology (ASCP) and Choosing Wisely

The American Board of Internal Medicine (ABIM) Foundation oversees the Choosing Wisely initiative where various professional societies can publish recommendations. The American Society for Clinical Pathology (ASCP) released a series of recommendations via Choosing Wisely beginning in 2013. In 2016, the ASCP released a recommendation clearly stating, "Do not test for amylase in cases of suspected acute pancreatitis. Instead, test for lipase.... Current guidelines and recommendations indicate that lipase should be preferred over total and pancreatic amylase for the initial diagnosis of acute pancreatitis and that the assessment should not be repeated over time to monitor disease prognosis." The ASCP also states that performing both lipase and amylase tests are not cost-effective given "marginally improving diagnostic efficiency compared to either marker alone." The ASCP recommendation does not mention any trypsin- or trypsinogen-based methodologies.

In 2020, the ASCP, along with Choosing Wisely and the ABIM Foundation, published a brochure titled *Thirty Things Physicians and Patients Should Question*. This brochure includes the following recommendation:

"Do not test for amylase in cases of suspected acute pancreatitis. Instead, test for lipase.

Amylase and lipase are digestive enzymes normally released from the acinar cells of the exocrine pancreas into the duodenum. Following injury to the pancreas, these enzymes are released into the circulation. While amylase is cleared in the urine, lipase is reabsorbed back into the circulation. In cases of acute pancreatitis, serum activity for both enzymes are greatly increased.

Serum lipase is now the preferred test due to its improved sensitivity, particularly in alcohol-induced pancreatitis. Its prolonged elevation creates a wider diagnostic window than amylase. In acute pancreatitis, amylase can rise rapidly within 3–6 hours of the onset of symptoms and may remain elevated for up to five days. Lipase, however, usually peaks at 24 hours with serum concentrations remaining elevated for 8–14 days. This means it is far more useful than amylase when the clinical presentation or testing has been delayed for more than 24 hours.

Current guidelines and recommendations indicate that lipase should be preferred over total and pancreatic amylase for the initial diagnosis of acute pancreatitis and that the assessment should not be repeated over time to monitor disease prognosis. Repeat testing should be considered only when the patient has signs and symptoms of persisting pancreatic or peripancreatic inflammation, blockage of the pancreatic duct or development of a pseudocyst. Testing both amylase and lipase is generally discouraged because it increases costs while only marginally improving diagnostic efficiency compared to either marker alone" (ASCP, 2020).

North American Society for Pediatric Gastroenterology, Hepatology and Nutrition Pancreas Committee (NASPGHAN)

NASPGHAN states that the primary biomarkers used to diagnose AP are serum lipase and amylase and note that "a serum lipase or amylase level of at least 3 times the upper limit of normal is considered consistent with pancreatitis." Further, NASPGHAN acknowledges that other biomarkers for diagnosis and management of AP have been investigated, but none are prominent and "many have yet to be validated for general clinical use" (NASPGHAN, 2018).





American Pancreatic Association (APA)

The APA acknowledges the utility of the measure of serum lipase and amylase for assessing pancreatic function "especially if they have a low value" (Conwell et al., 2014).

All societies reviewed prefer the use of serum lipase over serum amylase in the diagnosis of AP based on the higher sensitivity and selectivity of lipase. No consensus concerning the diagnostic threshold is reached between all of the societies where some use a threshold based on the Atlanta classification system, some do not specify a threshold, and one consortium recommends a time-based value system. A table adapted from the review by Lippi et al. (2012) summarizing the recommendations from various societies is below. This table has also been modified and updated with recent guidelines.

Table 2. Synthesis of available guidelines and recommendations for laboratory testing in the diagnosis of acute pancreatitis

Organization(s)	Preferred biomarker	Diagnostic threshold
Société Nationale Française de Gastro- Entérologie	Lipase	≥ 3 times the URL [upper limit of reference]
Japanese Society of Emergency Abdominal Medicine	Lipase	Not set
British Society of Gastroenterology; Association of Surgeons of Great Britain and Ireland, Pancreatic Society of Great Britain and Ireland, Association of Upper GI Surgeons of Great Britain and Ireland	Lipase	Value interpreted according to the time since the onset of symptoms
American of Gastroenterology	Lipase	\geq 2 to \geq 4 times the URL
American Gastroenterological Association	Lipase	≥ 3 times the URL
American of Family Physicians	Lipase	Not set
Japanese Ministry of Health, Labour, and Welfare	Lipase	Not set
Working Group of the Italian Association for the Study of the Pancreas	Lipase	Not set
World Society of Emergency Surgery*	Lipase	3 times the URL
North American Society for Pediatric Gastroenterology, Hepatology and Nutrition Pancreas Committee*	Amylase or Lipase	3 times the URL
American Board of Internal Medicine (ABIM), American Society for Clinical Pathology (ASCP) and Choosing Wisely*	Lipase	Not stated





UK Working Party on Acute Pancreatitis*	Lipase	should not rely on arbitrary limits of values 3 or 4 times greater than normal
American College of Gastroenterology*	Amylase or Lipase	≥ 3 times the URL
International Association of Pancreatology and the American Pancreatic Association*	Amylase or Lipase	≥ 3 times the URL

* This information was not part of the original table and was added to reference additional guidelines

The World Society of Emergency Surgery (WSES)

The WSES published guidelines in 2019 on the management of severe AP. These guidelines were determined by a panel of experts in 2018. The following statement was issued: "The cut-off value of serum amylase and lipase is normally defined to be three times the upper limit." Further, it is noted that "Collectively, serum lipase is considered a more reliable diagnostic marker of AP than serum amylase. No single test shows optimal diagnostic accuracy, but most current guidelines and recommendations indicate that lipase should be preferred over total and p-amylase. The main reasons supporting lipase over both types of amylases for the diagnosis of acute pancreatitis include higher sensitivity and larger diagnostic window" (Leppaniemi et al., 2019).

American Psychiatric Association (APA)

The APA published guidelines in 2006 for the treatment of patients with eating disorders. As shown below, these guidelines mention pancreatitis in several different tables throughout the document (table 5, table 6 and table 7).

- Table 5. Physical complications of Anorexia Nervosa
 - Regarding gastrointestinal laboratory test results, the APA (2006) states the following: "Liver function and associated tests: Occasionally abnormal liver function test results, increased serum amylase in purging patients; abnormal results rarely due to pancreatitis"
- Table 6: Physical complications of Bulimia Nervosa
 - Regarding gastrointestinal signs, the APA (2006) states the following: "Enlarged salivary glands, occasional blood-streaked vomitus; in vomiters, possibly gastritis, esophagitis, gastroesophageal erosions, esophageal dysmotility patterns (including gastroesophageal reflux and, rarely, Mallory-Weiss [esophageal] or gastric tears), increased rates of pancreatitis; in chronic laxative abusers, possibly colonic dysmotility or melanosis coli"
 - Regarding gastrointestinal laboratory test results, the APA (2006) states the following: "Serum amylase: Increased serum amylase (if fractionation is available, usually salivary gland isoenzymes); increased pancreatic amylase (rare), possibly indicating laxative abuse or other causes for pancreatic inflammation or pancreatitis"
- Table 7: Laboratory Assessments for Patients with Eating Disorders
 - Regarding nonroutine assessments, the APA (2006) has included "Serum amylase (fractionated for salivary gland isoenzyme if available to rule out pancreatic involvement)" (APA, 2006).





Academy for Eating Disorders (AED) Medical Care Standards Committee

The AED has published a guide to medical care for eating disorders. A table is included in these guidelines which is titled *Diagnostic Tests Indicated for All Patients with A Suspected ED* [eating disorder]. In a subcategory, titled *Additional Diagnostic Tests to Consider*, these guidelines mention pancreatic enzymes (amylase and lipase), stating that amylase could increase due to vomiting and pancreatitis, and lipase could increase due to pancreatitis (AED, 2016).

The Canadian Journal of Surgery

The Canadian Journal of Surgery released clinical practice guidelines on the management of acute pancreatitis. The following recommendations were made (Greenberg et al., 2016):

Guideline Recommendation		Strength of Evidence
1.	"A serum lipase test should be performed in all patients with a suspected diagnosis of acute pancreatitis. A 3-fold elevation of serum lipase from the upper limit of normal is required to make the diagnosis of acute pancreatitis	Strong
2.	C-reactive protein should be assessed at admission and daily for the first 72 hours after admission	Weak
3.	A serum C-reactive protein (CRP) level of 14 286 nmol/L (150 mg/dL) or greater at baseline or in the first 72 hours is suggestive of severe acute pancreatitis and is predictive of a worse clinical course. Thus, CRP should be assessed at admission and daily for the first 72 hours after admission."	Strong

The American Association for Clinical Chemistry

The American Association for Clinical Chemistry released recommendations for amylase testing in diagnosis and management of acute pancreatitis. The AACC provides the following recommendations:

- 1. "For diagnosis and management of acute pancreatitis, do not order the amylase test if serum lipase test is available.
- 2. The amylase test is not specific for pancreatitis and may be elevated due to other, non-pancreatic causes (such as acute cholecystitis, inflammatory bowel disease, intestinal obstruction, certain cancers, salivary disease, macroamylasemia, etc.)
- 3. Consider ordering this test when serum lipase is not available as a stat test and the patient presents with a sudden onset of abdominal pain with nausea and vomiting, fever, hypotension, and abdominal distension.
- 4. Testing both amylase and lipase should be discouraged because it increases costs while only marginally improving diagnostic efficiency compared to lipase alone" (AACC, 2020).

Chinese Pancreatic Surgery Association, Chinese Society of Surgery, Chinese Medical Association

The following associations discusses guidelines for the diagnosis of acute pancreatitis in China. "The diagnostic criteria of AP include (1) persistent upper abdominal pain, (2) concentration of serum amylase





and/or lipase at least 3 times higher than the normal upper limit, and (3) abdominal imaging findings suggestive of AP. AP can be diagnosed if 2 of the above 3 criteria are met." The guidelines also state that "Laboratory examination results show elevated serum amylase and lipase levels in patients with AP, and lipase has a higher specificity for the diagnosis of AP than amylase. Elevation of serum amylase and lipase does not affect the severity of AP" (Zhi, 2021).

VII. Applicable State and Federal Regulations

DISCLAIMER: If there is a conflict between this Policy and any relevant, applicable government policy for a particular member [e.g., Local Coverage Determinations (LCDs) or National Coverage Determinations (NCDs) for Medicare and/or state coverage for Medicaid], then the government policy will be used to make the determination. For the most up-to-date Medicare policies and coverage, please visit the Medicare search website: <u>http://www.cms.gov/medicare-coverage-database/overview-and-quick-search.aspx</u>. For the most up-to-date Medicaid policies and coverage, visit the applicable state Medicaid website.

Food and Drug Administration (FDA)

Many labs have developed specific tests that they must validate and perform in house. These laboratorydeveloped tests (LDTs) are regulated by the Centers for Medicare and Medicaid (CMS) as highcomplexity tests under the Clinical Laboratory Improvement Amendments of 1988 (CLIA '88). LDTs are not approved or cleared by the U. S. Food and Drug Administration; however, FDA clearance or approval is not currently required for clinical use.

Amylase

The FDA has approved multiple tests for human serum total amylase as well as for pancreatic amylase.

Lipase

The FDA has approved multiple tests for human serum lipase.

Trypsinogen/Trypsin/TAP

Trypsin immunostaining, trypsinogen-2 dipstick, and TAP serum tests are considered laboratory developed tests (LDT); developed, validated and performed by individual laboratories.

LDTs are regulated by the Centers for Medicare and Medicaid (CMS) as high-complexity tests under the Clinical Laboratory Improvement Amendments of 1988 (CLIA'88).

As an LDT, the U. S. Food and Drug Administration has not approved or cleared this test; however, FDA clearance or approval is not currently required for clinical use.

CRP

The FDA has approved multiple tests for human CRP, including assays for conventional CRP, high sensitivity CRP (hsCRP), and cardiac CRP (cCRP). On September 22, 2005, the FDA issued guidelines concerning the assessment of CRP (FDA, 2005).

Procalcitonin





On April 18, 2017, the FDA approved the Diazyme Procalcitonin PCT Assay, Diazyme Procalcitonin Calibrator Set, and Diazyme Procalcitonin Control Set as substantially equivalent and has received FDA 510K clearance for marketing.

On July 16, 2018, the FDA approved the Atellica IM BRAHMS Procalcitonin (PCT) device which is able to detect and measure procalcitonin (Pct) in human clinical specimens (FDA, 2018).

IL-6/IL-8

IL-6 and IL-8 are ELISA-based tests and are considered laboratory developed tests (LDT); developed, validated and performed by individual laboratories. IL-6 and IL-8 can also be components of a cytokine panel test, which is also an LDT.

VIII. Applicable CPT/HCPCS Procedure Codes

Procedure codes appearing in medical policy documents are only included as a general reference. This list may not be all inclusive and is subject to updates. In addition, codes listed are not a guarantee of payment.

Code	
Number	Code Description
82150	Amylase
83519	Immunoassay for analyte other than infectious agent antibody or infectious agent antigen; quantitative, by radioimmunoassay (eg, RIA)
00010	Immunoassay for analyte other than infectious agent antibody or infectious agent
83520	antigen; quantitative, not otherwise specified
83529	Interleukin-6 (IL-6)
83690	Lipase
84145	Procalcitonin (PCT)
86140	C-reactive protein

Current Procedural Terminology[©] American Medical Association. All Rights reserved.

IX. Evidence-based Scientific References

- AACC. (2020). AACC's Guide to Lab Test Utilization Retrieved from <u>https://www.aacc.org/advocacy-and-outreach/optimal-testing-guide-to-lab-test-utilization/a-f/amylase</u>
- AED. (2016). EATING DISORDERS A GUIDE TO MEDICAL CARE. Retrieved from <u>http://www.nyeatingdisorders.org/pdf/AED%20Medical%20Management%20Guide%203rd%20</u> <u>Edition.pdf</u>
- APA. (2006). Treatment of patients with eating disorders, third edition. American Psychiatric Association.
 Am J Psychiatry, 163(7 Suppl), 4-54. Retrieved from
 https://psychiatryonline.org/pb/assets/raw/sitewide/practice_guidelines/guidelines/eatingdisor_ders.pdf
- ASCP. (2020). Thirty Things Physicians and Patients Should Question. Retrieved from <u>https://www.ascp.org/content/docs/default-source/get-involved-</u> pdfs/istp_choosingwisely/2019_ascp-30-things-list.pdf





- Banks, P., & Freeman, M. (2006). Practice Guidelines in Acute Pancreatitis. *The American Journal Of Gastroenterology*, *101*, 2379. doi:10.1111/j.1572-0241.2006.00856.x
- Banks, P. A., Bollen, T. L., Dervenis, C., Gooszen, H. G., Johnson, C. D., Sarr, M. G., . . . Vege, S. S. (2013). Classification of acute pancreatitis—2012: revision of the Atlanta classification and definitions by international consensus. *Gut, 62*, 102-111.
- Basnayake, C., & Ratnam, D. (2015). Blood tests for acute pancreatitis. *Australian Prescriber, 38*(4), 128-130. doi:10.18773/austprescr.2015.043
- Bollen, T. L., Hazewinkel, M., & Smithuis, R. (2015, 05/01/2015). Pancreas-Acute Pancreatitis 2.0 2012 Revised Atlanta Classification of Acute Pancreatitis. *Radiology Assistant*. Retrieved from http://www.radiologyassistant.nl/en/p550455dae5806/pancreas-acute-pancreatitis-20.html
- Borowitz, D., Grant, R., & Durie, P. (1995, 2016). Pancreatic Enzymes Clinical Care Guidelines. Retrieved from <u>https://www.cff.org/Care/Clinical-Care-Guidelines/Nutrition-and-GI-Clinical-Care-Guidelines/Pancreatic-Enzymes-Clinical-Care-Guidelines/</u>
- Bradley, E. L., & Iii. (1993). A clinically based classification system for acute pancreatitis: Summary of the international symposium on acute pancreatitis, atlanta, ga, september 11 through 13, 1992.
 Archives of Surgery, 128(5), 586-590. doi:10.1001/archsurg.1993.01420170122019
- Burkart, J., Haigler, S., Caruana, R., & Hylander, B. (1991). Usefulness of peritoneal fluid amylase levels in the differential diagnosis of peritonitis in peritoneal dialysis patients. *J Am Soc Nephrol*, 1(10), 1186-1190.
- Ceylan, M. E., Evrensel, A., & Önen Ünsalver, B. (2016). Hyperamylasemia Related to Sertraline. *Korean Journal of Family Medicine*, *37*(4), 259-259. doi:10.4082/kjfm.2016.37.4.259
- Clavien, P. A., Robert, J., Meyer, P., Borst, F., Hauser, H., Herrmann, F., . . . Rohner, A. (1989). Acute pancreatitis and normoamylasemia. Not an uncommon combination. *Ann Surg*, *210*(5), 614-620.
- Coffey, M. J., Nightingale, S., & Ooi, C. Y. (2013). Serum Lipase as an Early Predictor of Severity in Pediatric Acute Pancreatitis. *Journal of Pediatric Gastroenterology and Nutrition, 56*(6), 602-608. doi:10.1097/MPG.0b013e31828b36d8
- Conwell, D. L., Lee, L. S., Yadav, D., Longnecker, D. S., Miller, F. H., Mortele, K. J., . . . Vege, S. S. (2014). American Pancreatic Association Practice Guidelines in Chronic Pancreatitis: Evidence-Based Report on Diagnostic Guidelines. *Pancreas*, *43*(8). Retrieved from <u>https://journals.lww.com/pancreasjournal/Fulltext/2014/11000/American Pancreatic Associati</u> <u>on Practice.6.aspx</u>
- Crockett, S. D., Wani, S., Gardner, T. B., Falck-Ytter, Y., & Barkun, A. N. (2018). American Gastroenterological Association Institute Guideline on Initial Management of Acute Pancreatitis. *Gastroenterology*, 154(4), 1096-1101. doi:10.1053/j.gastro.2018.01.032
- Eastler, J. (2017, 08/14/2017). Urine Trypsinogen 2 Dipstick for the Early Detection of Post-ERCP Pancreatitis. Retrieved from

https://clinicaltrials.gov/ct2/show/record/NCT03098082?view=record

- El Halabi, M., Bou Daher, H., Rustom, L. B. O., Marrache, M., Ichkhanian, Y., Kahil, K., . . . Sharara, A. I. (2019). Clinical utility and economic burden of routine serum lipase determination in the Emergency Department. *International Journal of Clinical Practice*, *73*(12), e13409. doi:<u>https://doi.org/10.1111/ijcp.13409</u>
- FDA. (2005). Review Criteria for Assessment of C-Reactive Protein (CRP), High Sensitivity C-Reactive Protein (hsCRP)k and Cardiac C-Reactive Protein (cCRP) Assays. Rockville, MD: U.S. Department of Health and Human Services Retrieved from <u>https://www.fda.gov/downloads/medicaldevices/deviceregulationandguidance/guidancedocum</u> ents/ucm071017.pdf
- FDA. (2018). Atellica IM BRAHMS Procalcitonin (PCT). Retrieved from <u>https://www.accessdata.fda.gov/scripts/cdrh/devicesatfda/index.cfm?db=pmn&id=K181002</u>





- Freedman, S. D. (2020). Clinical manifestations and diagnosis of chronic pancreatitis in adults. *UpToDate*. Retrieved from <u>https://www.uptodate.com/contents/clinical-manifestations-and-diagnosis-of-chronic-pancreatitis-in-adults</u>
- Furey, C., Buxbaum, J., & Chambliss, A. B. (2020). A review of biomarker utilization in the diagnosis and management of acute pancreatitis reveals amylase ordering is favored in patients requiring laparoscopic cholecystectomy. *Clin Biochem*, 77, 54-56. doi:10.1016/j.clinbiochem.2019.12.014
- Greenberg, J. A., Hsu, J., Bawazeer, M., Marshall, J., Friedrich, J. O., Nathens, A., . . . McLeod, R. S. (2016). Clinical practice guideline: management of acute pancreatitis. *Canadian journal of surgery. Journal canadien de chirurgie, 59*(2), 128-140. doi:10.1503/cjs.015015
- Guidelines, W. G. I. A. A. P. (2013). IAP/APA evidence-based guidelines for the management of acute pancreatitis. *Pancreatology*, *13*(4), e1-e15. doi:10.1016/j.pan.2013.07.063
- Gumaste, V. V., Dave, P. B., Weissman, D., & Messer, J. (1991). Lipase/amylase ratio. A new index that distinguishes acute episodes of alcoholic from nonalcoholic acute pancreatitis. *Gastroenterology*, *101*(5), 1361-1366.
- Hagjer, S., & Kumar, N. (2018). Evaluation of the BISAP scoring system in prognostication of acute pancreatitis - A prospective observational study. *Int J Surg*, *54*(Pt A), 76-81. doi:10.1016/j.ijsu.2018.04.026
- Herrmann-Storck, C., Saint Louis, M., Foucand, T., Lamaury, I., Deloumeaux, J., Baranton, G., . . . Cornet, M. (2010). Severe Leptospirosis in Hospitalized Patients, Guadeloupe. *Emerging Infectious Diseases, 16*(2), 331-334. doi:10.3201/eid1602.090139
- Ismail, O. Z., & Bhayana, V. (2017). Lipase or amylase for the diagnosis of acute pancreatitis? *Clin Biochem*, *50*(18), 1275-1280. doi:10.1016/j.clinbiochem.2017.07.003
- Jakkampudi, A., Jangala, R., Reddy, R., Mitnala, S., Rao, G. V., Pradeep, R., . . . Talukdar, R. (2017). Acinar injury and early cytokine response in human acute biliary pancreatitis. *Sci Rep*, 7(1), 15276. doi:10.1038/s41598-017-15479-2
- Judal. (2022). Urinary amylase levels in the diagnosis of acute pancreatitis: a prospective case control study. *International Surgery Journal*. Retrieved from
 - https://www.ijsurgery.com/index.php/isj/article/view/8530
- Kemppainen, E. A., Hedstrom, J. I., Puolakkainen, P. A., Sainio, V. S., Haapiainen, R. K., Perhoniemi, V., . .
 Stenman, U. H. (1997). Rapid measurement of urinary trypsinogen-2 as a screening test for acute pancreatitis. *N Engl J Med*, *336*(25), 1788-1793. doi:10.1056/nejm199706193362504
- Khanna, A. K., Meher, S., Prakash, S., Tiwary, S. K., Singh, U., Srivastava, A., & Dixit, V. K. (2013). Comparison of Ranson, Glasgow, MOSS, SIRS, BISAP, APACHE-II, CTSI Scores, IL-6, CRP, and Procalcitonin in Predicting Severity, Organ Failure, Pancreatic Necrosis, and Mortality in Acute Pancreatitis. *HPB Surgery, 2013*, 367581. doi:10.1155/2013/367581
- Kumar, P., Ghosh, A., Tandon, V., & Sahoo, R. (2016). Gullo's Syndrome: A Case Report. *Journal of Clinical and Diagnostic Research : JCDR, 10*(2), OD21-OD22. doi:10.7860/JCDR/2016/17038.7285
- Lempinen, M., Kylänpää-Bäck, M.-L., Stenman, U.-H., Puolakkainen, P., Haapiainen, R., Finne, P., . . . Kemppainen, E. (2001). Predicting the Severity of Acute Pancreatitis by Rapid Measurement of Trypsinogen-2 in Urine. *Clinical Chemistry*, 47(12), 2103. Retrieved from <u>http://clinchem.aaccjnls.org/content/47/12/2103.abstract</u>
- Leppaniemi, A., Tolonen, M., Tarasconi, A., Segovia-Lohse, H., Gamberini, E., Kirkpatrick, A. W., . . . Catena, F. (2019). 2019 WSES guidelines for the management of severe acute pancreatitis. *World J Emerg Surg, 14*, 27. doi:10.1186/s13017-019-0247-0
- Levy, P., Boruchowicz, A., Hastier, P., Pariente, A., Thevenot, T., Frossard, J. L., . . . Ruszniewski, P. (2005). Diagnostic criteria in predicting a biliary origin of acute pancreatitis in the era of endoscopic ultrasound: multicentre prospective evaluation of 213 patients. *Pancreatology*, 5(4-5), 450-456. doi:10.1159/000086547





- Li, N., Wang, B. M., Cai, S., & Liu, P. L. (2018). The Role of Serum High Mobility Group Box 1 and Interleukin-6 Levels in Acute Pancreatitis: A Meta-Analysis. *J Cell Biochem*, *119*(1), 616-624. doi:10.1002/jcb.26222
- Lippi, G., Valentino, M., & Cervellin, G. (2012). Laboratory diagnosis of acute pancreatitis: in search of the Holy Grail. *Critical Reviews in Clinical Laboratory Sciences*, 49(1), 18-31. doi:10.3109/10408363.2012.658354
- Liu, P., Xiao, Z., Yan, H., Lu, X., Zhang, X., Luo, L., . . . Zhu, Y. (2021). Serum Amylase and Lipase for the Prediction of Pancreatic Injury in Critically III Children Admitted to the PICU. *Pediatric Critical Care Medicine*, *22*(1), e10-e18. doi:10.1097/pcc.00000000002525
- Liu, S., Wang, Q., Zhou, R., Li, C., Hu, D., Xue, W., . . . Peng, A. (2016). Hyperamylasemia as an Early Predictor of Mortality in Patients with Acute Paraquat Poisoning. *Medical Science Monitor : International Medical Journal of Experimental and Clinical Research, 22*, 1342-1348. doi:10.12659/MSM.897930
- NASPGHAN. (2018). Management of Acute Pancreatitis in the Pediatric Population: A Clinical Report From the North American Society for Pediatric Gastroenterology, Hepatology and Nutrition Pancreas Committee. Retrieved from

https://www.naspghan.org/files/Management_of_Acute_Pancreatitis_in_the_Pediatric.33.pdf

- Neoptolemos, J. P., Kemppainen, E. A., Mayer, J. M., Fitzpatrick, J. M., Raraty, M. G., Slavin, J., . . . Puolakkainen, P. A. (2000). Early prediction of severity in acute pancreatitis by urinary trypsinogen activation peptide: a multicentre study. *Lancet*, *355*(9219), 1955-1960.
- Pacheco, R. C., & Oliveira, L. C. (2007). [Lipase/amylase ratio in biliary acute pancreatitis and alcoholic acute/acutized chronic pancreatitis]. *Arq Gastroenterol, 44*(1), 35-38.
- Pezzilli, R., Venturi, M., Morselli-Labate, A. M., Ceciliato, R., Lamparelli, M. G., Rossi, A., . . . Corinaldesi, R. (2004). Serum Trypsinogen Activation Peptide in the Assessment of the Diagnosis and Severity of Acute Pancreatic Damage: A Pilot Study Using a New Determination Technique. *Pancreas,* 29(4), 298-305. Retrieved from

https://journals.lww.com/pancreasjournal/Fulltext/2004/11000/Serum_Trypsinogen_Activation Peptide in the.9.aspx

- Quinlan, J. D. (2014). Acute pancreatitis. Am Fam Physician, 90(9), 632-639.
- Rau, B. M., Kemppainen, E. A., Gumbs, A. A., Buchler, M. W., Wegscheider, K., Bassi, C., . . . Beger, H. G. (2007). Early assessment of pancreatic infections and overall prognosis in severe acute pancreatitis by procalcitonin (PCT): a prospective international multicenter study. *Ann Surg*, 245(5), 745-754. doi:10.1097/01.sla.0000252443.22360.46
- Rompianesi, G., Hann, A., Komolafe, O., Pereira, S. P., Davidson, B. R., & Gurusamy, K. S. (2017). Serum amylase and lipase and urinary trypsinogen and amylase for diagnosis of acute pancreatitis. *Cochrane Database of Systematic Reviews*(4). doi:10.1002/14651858.CD012010.pub2
- Sendler, M., & Lerch, M. M. (2020). The Complex Role of Trypsin in Pancreatitis. *Gastroenterology*, *158*(4), 822-826. doi:10.1053/j.gastro.2019.12.025
- Simha, A., Saroch, A., Pannu, A. K., Dhibar, D. P., Sharma, N., Singh, H., & Sharma, V. (2021). Utility of point-of-care urine trypsinogen dipstick test for diagnosing acute pancreatitis in an emergency unit. *Biomarkers in Medicine*, 15(14), 1271-1276. doi:10.2217/bmm-2021-0067
- Simsek, O., Kocael, A., Kocael, P., Orhan, A., Cengiz, M., Balci, H., . . . Uzun, H. (2018). Inflammatory mediators in the diagnosis and treatment of acute pancreatitis: pentraxin-3, procalcitonin and myeloperoxidase. *Arch Med Sci*, *14*(2), 288-296. doi:10.5114/aoms.2016.57886
- Stevens, T., & Conwell, D. L. (2020, 10/05/2016). Exocrine pancreatic insufficiency. *UpToDate*. Retrieved from https://www.uptodate.com/contents/exocrine-pancreatic-insufficiency





- Tenner, S., Baillie, J., DeWitt, J., Vege, S. S., & American College of, G. (2013). American College of Gastroenterology guideline: management of acute pancreatitis. *Am J Gastroenterol, 108*(9), 1400-1415; 1416. doi:10.1038/ajg.2013.218
- Tenner, S. M., & Steinberg, W. (1992). The admission serum lipase:amylase ratio differentiates alcoholic from nonalcoholic acute pancreatitis. *Am J Gastroenterol, 87*(12), 1755-1758.
- Terui, K., Hishiki, T., Saito, T., Mitsunaga, T., Nakata, M., & Yoshida, H. (2013). Urinary amylase / urinary creatinine ratio (uAm/uCr) - a less-invasive parameter for management of hyperamylasemia. BMC Pediatrics, 13, 205-205. doi:10.1186/1471-2431-13-205
- Tian, F., Li, H., Wang, L., Li, B., Aibibula, M., Zhao, H., . . . Ma, X. (2020). The diagnostic value of serum C-reactive protein, procalcitonin, interleukin-6 and lactate dehydrogenase in patients with severe acute pancreatitis. *Clinica Chimica Acta*, *510*, 665-670. doi:https://doi.org/10.1016/j.cca.2020.08.029
- Toouli, J., Brooke-Smith, M., Bassi, C., Carr-Locke, D., Telford, J., Freeny, P., . . . Tandon, R. (2002). Guidelines for the management of acute pancreatitis. *J Gastroenterol Hepatol*, *17 Suppl*, S15-39.
- Vege, S. S. (2020, 05/21/2018). Approach to the patient with elevated serum amylase or lipase. *UpToDate*. Retrieved from <u>https://www.uptodate.com/contents/approach-to-the-patient-with-elevated-serum-amylase-or-lipase</u>
- Vege, S. S. (2021a). Clinical manifestations and diagnosis of acute pancreatitis. *UpToDate*. Retrieved from <u>https://www.uptodate.com/contents/clinical-manifestations-and-diagnosis-of-acute-pancreatitis</u>
- Vege, S. S. (2021b, 02/24/2017). Pathogenesis of acute pancreatitis. *UpToDate*. Retrieved from <u>https://www.uptodate.com/contents/pathogenesis-of-acute-pancreatitis</u>
- Ventrucci, M., Pezzilli, R., Naldoni, P., Plate, L., Baldoni, F., Gullo, L., & Barbara, L. (1987). Serum pancreatic enzyme behavior during the course of acute pancreatitis. *Pancreas, 2*(5), 506-509.
- Wei, M., Xie, X., Yu, X., Lu, Y., Ke, L., Ye, B., . . . Li, J. (2022). Predictive value of serum cholinesterase in the mortality of acute pancreatitis: A retrospective cohort study. *European Journal of Clinical Investigation, n/a*(n/a), e13741. doi:<u>https://doi.org/10.1111/eci.13741</u>
- Wilson, J., & Smith, R. (2015). Chronic pancreatitis and the use of PERT. In A. P. C. P. G. W. Party (Ed.), Australasian guidelines for the management of pancreatic exocrine insufficiency (Vol. Oct 2015). Ourimbah, NSW, Australia: Australasian Pancreatic Club.
- Witt, H., Apte, M. V., Keim, V., & Wilson, J. S. (2007). Chronic pancreatitis: challenges and advances in pathogenesis, genetics, diagnosis, and therapy. *Gastroenterology*, 132(4), 1557-1573. doi:10.1053/j.gastro.2007.03.001
- Wolfe, B. E., Jimerson, D. C., Smith, A., & Keel, P. K. (2011). Serum Amylase in Bulimia Nervosa and Purging Disorder: Differentiating the Association with Binge Eating versus Purging Behavior. *Physiology & behavior*, 104(5), 684-686. doi:10.1016/j.physbeh.2011.06.025
- Yadav, D., Agarwal, N., & Pitchumoni, C. S. (2002). A critical evaluation of laboratory tests in acute pancreatitis. *Am J Gastroenterol, 97*(6), 1309-1318. doi:10.1111/j.1572-0241.2002.05766.x
- Yang, H., Wang, H., Chavan, S. S., & Andersson, U. (2015). High Mobility Group Box Protein 1 (HMGB1): The Prototypical Endogenous Danger Molecule. *Mol Med, 21 Suppl 1*, S6-s12. doi:10.2119/molmed.2015.00087
- Yasuda, H., Kataoka, K., Takeyama, Y., Takeda, K., Ito, T., Mayumi, T., . . . Shimosegawa, T. (2019).
 Usefulness of urinary trypsinogen-2 and trypsinogen activation peptide in acute pancreatitis: A multicenter study in Japan. *World J Gastroenterol, 25*(1), 107-117. doi:10.3748/wjg.v25.i1.107
- Zhan, X., Wan, J., Zhang, G., Song, L., Gui, F., Zhang, Y., . . . Ji, B. (2019). Elevated intracellular trypsin exacerbates acute pancreatitis and chronic pancreatitis in mice. Am J Physiol Gastrointest Liver Physiol, 316(6), G816-g825. doi:10.1152/ajpgi.00004.2019





Zhi, Z. W. K. Z. (2021). [Guidelines for diagnosis and treatment of acute pancreatitis in China (2021)]. *Zhonghua Wai Ke Za Zhi, 59*(7), 578-587. doi:10.3760/cma.j.cn112139-20210416-00172

X. Revision History

Revision Date	Summary of Changes	
12/01/2021	Initial Effective Date	
04/12/2022	Updated background, guidelines, and evidence-based scientific references. Literature review did not necessitate any modifications to the coverage criteria. Code number 83529 added to Procedure Codes. Added statement "Procedure codes appearing in Medical Policy documents are included only as a general reference tool for each policy. They may not be all-inclusive" at end of procedure code section.	
10/02/2022	Updated background, guidelines, and evidence-based scientific references. Literature review necessitated the following changes to coverage criteria: CC1 and CC2, addition of "(preferred)" after serum lipase Addition of "or urine" to CC3b, now reads: "b) serum or urine trypsin/trypsinogen/TAP (trypsinogen activation peptide)" Addition of new CC5: "Measurement of urinary amylase concentration for the initial diagnosis of acute pancreatitis DOES NOT MEET COVERAGE CRITERIA in all patients presenting with signs and symptoms of acute pancreatitis* (please see Note 1)" Revised code disclaimer statement	