Prophylactic ulinastatin administration for preventing post-endoscopic retrograde cholangiopancreatography pancreatitis: A meta-analysis

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Abstract. The objective of the present study was to perform a meta-analysis of all available studies on the effect of prophylactic ulinastatin administration on preventing post-endoscopic retrograde cholangiopancreatography (ERCP) pancreatitis (PEP). The PubMed, Web of Knowledge and Chinese National Knowledge Infrastructure databases were searched to identify all relevant studies published in English or Chinese prior to April 2016. Cochrane Review Manager was used to calculate the pooled risk ratio (RR) and 95% confidence interval (CI) to determine the effect of prophylactic ulinastatin on PEP, post-ERCP hyperamylasemia (PEHA) and post-ERCP abdominal pain. The analysis revealed that prophylactic ulinastatin administration significantly reduced the PEP risk (RR=0.49; 95% CI: 0.33-0.74; P=0.0006; I²=24); however, such significant risk reduction occurred only in patients with low or average risk for PEP and high-dosage ulinastatin (150,000 or 200,000 U) administration, and when the ulinastatin administration began prior to or during ERCP. Pre-ERCP ulinastatin administration alone without additional administration after ERCP was sufficient. Prophylactic ulinastatin also significantly reduced the PEHA risk (RR=0.68; 95% CI: 0.56-0.83; P=0.0001; I²=19) and marginally reduced the incidence of post-ERCP abdominal pain (RR=0.67; 95% CI: 0.45-1.00; P=0.05; I²=67). In conclusion, prophylactic ulinastatin administration significantly reduced the risk of PEP in patients with low or average risk for PEP when administered at a high dosage prior to or during ERCP. High-quality studies, particularly on high-risk patients, are warranted.

Introduction

Endoscopic retrograde cholangiopancreatography (ERCP) has been a widely accepted procedure for diagnosing and evaluating pancreatic and biliary diseases ever since its introduction in the 1970s, and it remains the first choice in treating and managing various pancreaticobiliary diseases (1,2). However, it also has the disadvantage of carrying the highest complication rate among all of the gastrointestinal endoscopic procedures currently in practice (1,2). Post-ERCP complications include pancreatitis, cholangitis, cholecystitis, perforation and bleeding; among them, post-ERCP pancreatitis (PEP) is most common. Its estimated incidence ranges from 1-10% in average-risk patients, and reaches nearly 30% in high-risk patients (1,2). Although PEP is mild or moderate in most of the cases, it is severe in ~10% of the cases and may potentially be fatal (3,4). Given the relatively high incidence of PEP and its potentially severe consequences, it is no surprise that the interest in developing pharmaceutical preventions for PEP has been growing (2). However, studies on the preventive effects of various pharmaceutical agents on PEP often produced inconsistent or even conflicting results and therefore, the 2010 European Society of Gastrointestinal Endoscopy (ESGE) guidelines for the prophylaxis of PEP stated that non-steroidal anti-inflammatory drugs are the only ones with proven efficacy (5).

Since proteolytic enzyme activation starting with activation of trypsinogen to trypsin in pancreatic acinar cells has a triggering and important role in the pathogenesis of PEP, protease inhibitors such as ulinastatin and gabexate mesylate are potential prophylactic agents for reducing the risk of PEP, as ulinastatin, a trypsin inhibitor, may prevent the initial phase of the pathogenesis of PEP (1,3,5). In fact, protease inhibitors such as ulinastatin and gabexate mesylate have already been approved by the Ministry of Health, Labor, and Welfare of Japan as therapeutic medications for acute pancreatitis, and they have been widely used in China and Japan to treat and manage acute pancreatitis (1,5). Compared to gabexate mesylate that has a 55-second plasma half-life, ulinastatin has a relatively long plasma half-life (35 min) and may be injected as a bolus as opposed to the 13-h continuous infusion required for gabexate mesylate to prevent pancreatic injury (1,3,6). Despite being commonly used in Japan for PEP prevention, numerous studies on the efficacy of prophylactic ulinastatin...
in preventing PEP have produced inconsistent results (4,6-17). Several meta-analyses on this topic also produced inconsistent results, with the study by Chen et al (3) published in 2010 finding ulinastatin effective in reducing the incidence of PEP and post-ERCP hyperamylasemia (PEHA), an early indication of possible PEP, while the study by Yuhara et al (1) published in 2014 concluded that ulinastatin was not associated with a reduced risk of PEP. Since then, several additional studies on the effect of ulinastatin in preventing PEP and PEHA have been published. Probably due to their inconsistent results, protease inhibitors such as ulinastatin were not recommended in the 2010 ESGE guidelines.

The present study performed a meta-analysis of all available relevant studies on the effect of prophylactic ulinastatin on preventing PEP, PEHA and also on post-ERCP abdominal pain, another early indication of possible PEP. The present analysis may provide further evidence as to whether ulinastatin may be used as a prophylactic medication for PEP.

Materials and methods

Literature search. The PubMed, Web of Knowledge and Chinese National Knowledge Infrastructure (CNKI) databases were systematically searched to identify all relevant studies published prior to April 2016 in English or Chinese. The following search terms were used: ‘post-endoscopic retrograde cholangiopancreatography’, ‘ERCP’, ‘pancreatitis’, ‘PEP’, ‘ulinastatin’, ‘urinary trypsin inhibitor’ and ‘UTI’. In addition, the reference lists of the identified studies and relevant reviews were examined to identify eligible studies possibly missed in the initial search.

Inclusion and exclusion criteria. All studies comparing the effect of prophylactic ulinastatin administration with a control in preventing PEP published in English or Chinese were considered eligible for the present meta-analysis. The inclusion criteria were as follows: i) The study was a controlled trial wherein patients were grouped into 2 treatment arms to receive either ulinastatin or a non-ulinastatin control (placebo or nothing), with all other treatments and medications being the same or comparable; ii) patients in the study received ERCP; and iii) the study reported on at least one of the following outcome measures: Incidence of PEP, PEHA or post-ERCP abdominal pain in the 2 groups.

Abstracts, reviews, editorials and studies without a proper control or with only one treatment arm were excluded.

KZ and JPW did a first round of screening by independently reviewing titles and abstracts of the studies identified by the initial search in order to shorten the list of relevant studies. They then reviewed the full-text studies on the shortened list to obtain a final list of studies to be included in the meta-analysis. Any disagreement between the two investigators was resolved by discussions among all of the contributing authors in order to reach a consensus.

Assessment of study quality. KZ and JPW independently assessed the quality of each included study according to Jadad et al (18) based on the following points: i) Whether the trial is randomized and if it is, whether the randomization is appropriate; ii) whether the study is double-blinded and if it is, whether the double-blinding is done appropriately; iii) whether there is a proper description of patient withdrawal or dropout, using a score of 0 or 1 for each question. Therefore, a study's possible total Jadad score were within the range of 0-5. In the present meta-analysis, studies with a Jadad score of ≥3 were defined as a high-quality studies, those with a Jadad score of 2 as average-quality studies and studies with a Jadad score <2 as low-quality studies.

Data extraction. KZ and JGS independently reviewed the full text of each study and extracted the following data: First author name, year of publication, study design, sample size (ulinastatin/control), male/female ratio, route and time of drug administration and main outcome measures. In addition, for the meta-analysis on the effect of ulinastatin on PEP, the definition of PEP, incidence of PEP and summary results were also extracted; for the meta-analysis on the effect on PEHA, definition of PEHA, incidence of PEHA and summary results were also extracted; and for the meta-analysis on the effect of ulinastatin on post-ERCP abdominal pain, the incidence of post-ERCP abdominal pain and summary results were also extracted. The Jadad score of each included study was also determined. Any disagreement between the two reviewers was resolved by discussions between them.

Statistical analysis. Cochrane Review Manager (Revman, version 5.3; the Cochrane Institute, London, UK) was used to perform the statistical analysis. Pooled risk ratio (RR) and 95% confidence interval (CI) were used as the effect size measurements for the analysis, and they were first calculated with a random-effects model, as such a model expects and assumes a true diversity in the results of the included studies and hence includes a between-studies variance into its calculation (19). The Z test was used to assess the statistical significance of the pooled RRs, and a 95% CI not including the value of 1 and P<0.05 were considered to indicate statistical significance. Inter-study heterogeneities were assessed using the χ² test based on Cochrane Q statistics and when a P value for the Q statistics was <0.10, the heterogeneity was statistically significant (20). The degree of inter-study heterogeneities was further quantified using the I² index, with a higher I² value representing a greater degree of heterogeneity (an I² value of ~25, ~50 and ~75% indicated low, moderate and high degree of heterogeneity, respectively) (21). If no inter-study heterogeneity was found using the random-effects model, pooled RR and 95% CI were re-calculated using a fixed-effects model.

Sub-group analyses were performed for each of the 3 outcome measures based on the following factors: i) Whether the ulinastatin administration began prior to, during or after ERCP; ii) for those studies in which ulinastatin administration began prior to ERCP, whether ulinastatin was administered only prior to or also after ERCP; iii) whether the patients were at high risk of PEP; and iv) whether low- or high-dosage ulinastatin was administered. In addition, as part of a sensitivity analysis, sub-group analysis was also performed based on the qualities of the included studies in order to determine the impact of low-quality studies on the results and the stability and robustness of the analysis results. Sensitivity analysis further included change of the statistical model (a fixed-effects vs. a random-effects model). Finally, publication bias in the analysis was determined using a funnel plot.
Results

Eligible studies and study characteristics. Fig. 1 presents a flow chart illustrating the literature search process of the present study and its results. A total of 75 potentially eligible studies were identified by searching the PubMed, Web of Science and CNKI databases. After excluding 19 duplicate studies, 56 studies were further examined to determine whether they meet the inclusion criteria and should be included in the present meta-analyses. A total of 43 studies were excluded, as they were reviews or meta-analyses, abstracts, irrelevant to the topic of the present study or did not provide any of the required data. Finally, 13 studies were included in the present meta-analyses, and their characteristics are described in Table I. Among these 13 studies, 11 studies (4,6-9,11-14,16,17) were included in the present meta-analysis on the effect of prophylactic ulinastatin on PEP prevention (4,6-9,11-14,16,17). Prophylactic ulinastatin led to a significantly reduced risk of PEP compared with the control (RR=0.49; 95% CI: 0.33-0.74; P=0.0006; I^2=24); however, such significant risk reduction only occurred when ulinastatin administration began prior to (RR=0.46; 95% CI: 0.26-0.83; P=0.010; I^2=32) or during ERCP (RR=0.37; 95% CI: 0.19-0.75; P=0.005; I^2=0), but not after ERCP (RR=0.64; 95% CI: 0.15-2.70; P=0.55; I^2=60; Fig. 2A). In addition, ulinastatin administration prior to ERCP without additional administration after ERCP appeared to be sufficient to significantly reduce the PEP risk (RR=0.37; 95% CI: 0.15-0.90; P=0.03; I^2=0; Fig. 2B).

Further sub-group analysis revealed that the significantly reduced PEP risk associated with ulinastatin vs. control only occurred in patients with a low or average risk for PEP (RR=0.49; 95% CI: 0.33-0.72; P=0.0003; I^2=13), but not in high-risk patients (RR=0.29; 95% CI: 0.01-6.37; P=0.43; I^2=87; Fig. 2C). The significant PEP risk reduction effect of ulinastatin was only observed with high-dosage ulinastatin administration (150,000 or 200,000 U; RR=0.49; 95% CI: 0.32-0.74; P=0.0008; I^2=19), but not with low-dosage ulinastatin (100,000 U; RR=0.44; 95% CI: 0.10-1.88; P=0.27; I^2=55; Fig. 2D).

As a part of the sensitivity analysis, the present study evaluated the impact of the qualities of the studies on the results by dividing the studies into high- or average-quality studies (Jadad score, ≥2) and low-quality studies (Jadad score, <2). The sub-group analysis revealed that the qualities of the
| Author, year            | Language                  | Study design                                      | Sample size | Male/female | Age ulinastatin/control (years) | Drug administration                                                                 | Main outcome measure(s)                                      | Jadad score | Refs. |
|-------------------------|---------------------------|--------------------------------------------------|-------------|-------------|-------------------------------|--------------------------------------------------------------------------------------|-------------------------------------------------------------|-------------|-------|
| Ohwada et al, 1997      | English                   | Prospective, placebo controlled study            | 22/46       | -           | -                             | Ulinastatin: 2-6 ml, contrast medium containing 50,000 U ulinastatin injected into the pancreatic duct during ERCP; Control: 2-6 ml of the same contrast medium administered in the same manner | Incidence of abdominal pain and change of amylase level post ERCP | 0           | (10)  |
| Gong et al, 2004        | Chinese (with an English abstract) | Retrospective, placebo-controlled study          | 68/62       | -           | -                             | Ulinastatin: 200,000 U ulinastatin dissolved in 250 ml saline solution administered intravenously before and also for 3 days after ERCP; Control: 250 ml saline solution administered in the same manner | Incidence of PEP                                           | 1           | (12)  |
| Song et al, 2005        | Chinese (with an English abstract) | Prospective, randomized, placebo-controlled study | 20/20       | 9/11:8/12   | 48.4/46.8                      | Ulinastatin: 200,000 U ulinastatin dissolved in 250 ml of saline solution administered intravenously 1 h before ERCP; Control: 250 ml saline solution administered in the same manner | Incidence of PEP and PEHA                                   | 1           | (11)  |
| Tsujino et al, 2005     | English                   | Multicenter, prospective, randomized, double-blind, placebo-controlled study | 204/202     | 113/91:131/71 | 65 (22-97)/65(22-92)*           | Ulinastatin: 150,000 U dissolved in 100 ml 0.9% saline solution administered by intravenous drip infusion immediately before ERCP for 10 min; Control: 100 ml 0.9% saline solution was given in the same manner | Incidence of PEP stratified by severity, PEHA and abdominal pain | 5           | (6)   |
| Author, year | Language | Study design | Male/female | Age (years) | Sample size | Drug administration | Main outcome measure | Jadad score | Drug outcome | Refs. |
|--------------|----------|--------------|-------------|-------------|-------------|----------------------|---------------------|-------------|-------------|-------|
| Yao et al., 2005 | Chinese | Prospective, randomized, controlled study | 64:62 | 36/26 | 64/62 | Ulinastatin: 100,000 U | Incidence of abdominal pain and serum amylase level before and after ERCP | 2 | (15) |
| Yoo et al., 2008 | English | Multicenter, prospective, double-blind, placebo controlled study | 119/108 | 64/59 | 64/59 | Ulinastatin: 100,000 U | Incidence of acute PEP | 5 | (8) |
| Wang et al., 2010 | Chinese | Prospective, randomized, placebo controlled study | 93/85 | - | - | Ulinastatin: 100,000 U | Incidence of PEP | 1 | (14) |
| Yoo et al., 2012 | English | Retrospective study | 229/658 | 128/101 | 128/101 | Ulinastatin: 150,000 U | Incidence of PEP | 0 | (4) |
### Table I. Continued.

| Author, year | Language | Study design                      | Sample size ulinastatin/ control (n) | Male/female (ulinastatin: control, n) | Age ulinastatin/ control (years) | Drug administration                                                                 | Main outcome measure(s) | Jadad score | (Refs.) |
|--------------|----------|-----------------------------------|-------------------------------------|---------------------------------------|----------------------------------|----------------------------------------------------------------------------------|------------------------|------------|---------|
| Xiong et al, 2013 | Chinese | Prospective, randomized, controlled study | 37/36 | 19/18:16/20 | 59.8/60.1 | Ulinastatin: In addition to routine therapy, 200,000 U ulinastatin dissolved in 250 ml 5% dextrose solution administered by intravenous drip infusion right after ERCP and also 24 and 48 h after ERCP; Control: Routine therapy. | Incidence of PEP and PEHA, and also change of serum amylase level | 1 | (13) |
| Park et al, 2014 | English | Prospective, randomized, placebo-controlled study | 53/53 low-risk 11/37 high-risk 42/16 | 31/22:29/24 | 59.4±16.8/60.5±16.2 | Ulinastatin: 150,000 U ulinastatin dissolved in 1,000 ml 5% dextrose solution intravenously administered from 2-4 h before ERCP to 6-8 h after ERCP; Control: 1,000 ml 5% dextrose solution administered in the same manner. | Incidence of PEP stratified by severity and low-and high-risk patients, PEHA and abdominal pain | 2 | (9) |
| Wang et al, 2014 | English | Prospective, randomized control study | 160/120 | - | - | Ulinastatin: 40 ml meglumine diatrizoate contrast medium containing 200,000 U ulinastatin; Control: 40 ml meglumine diatrizoate contrast medium. Both administered during ERCP. | Incidence of PEP in the 2 groups | 2 | (7) |
| Chen et al, 2015 | Chinese | Prospective, randomized, placebo-controlled study | 60/60 | 35/25:37/23 | 58.3/60.2 | Ulinastatin: 100,000 U ulinastatin dissolved in 250 ml 0.9% saline solution administered by intravenous drip infusion 30 min before, 1 h and 24 h after ERCP; Control: 250 ml 0.9% saline solution administered in the same manner. | Incidence of PEP, PEHA and abdominal pain | 1 | (16) |
Effect of prophylactic ulinastatin administration on preventing PEHA. As presented in Table III, 9 studies including 875 patients receiving prophylactic ulinastatin and 1,281 patients receiving a placebo or no drug as a control were included in the meta-analysis on the efficacy of prophylactic ulinastatin in preventing PEHA (4,6,8,9,11,13,14,16,17). The analysis revealed that prophylactic ulinastatin led to a significant PEHA risk reduction vs. the control (RR=0.68; 95% CI: 0.56-0.83; P=0.0001; I²=19); however, such significantly reduced PEHA risk only occurred when the ulinastatin administration began prior to ERCP (RR=0.62; 95% CI: 0.50-0.77; P<0.0001; I²=0), but not during or after ERCP (Fig. 3A). Furthermore, post-ERCP ulinastatin administration in addition to the pre-ERCP administration appeared to be necessary to significantly reduce the PEHA risk (RR=0.65; 95% CI: 0.51-0.84; P=0.00009; I²=0; Fig. 3B).

Furthermore, a significantly reduced PEHA risk associated with ulinastatin vs. control was observed in patients with a low or average risk for PEP (RR=0.66; 95% CI: 0.54-0.82; P=0.0001; I²=13), but not in high-risk patients (RR=0.71; 95% CI: 0.19-2.67; P=0.61; I²=89; Fig. 3C). Ulinastatin was able to significantly reduce the risk of PEHA if administered at a high dose (150,000 or 200,000 U; RR=0.70; 95% CI: 0.58-0.85; P=0.0004; I²=7), but not at a low dose (100,000 U; Fig. 3D).

Furthermore sub-group analysis based on the quality of the studies revealed that the quality of the studies had an impact on the results: Although analysis of low-quality studies indicated a significant reduction of the risk of PEHA associated with ulinastatin vs. control, analysis of high- or average-quality studies only revealed a trend short of statistically significance of the reduced PEHA risk associated with ulinastatin (Fig. 3E). In most analyses, using a fixed-effects model did not significantly impact the results obtained by using a random-effects model; however, there were a few exceptions. A fixed-effects model analysis indicated a statistically significant PEHA risk reduction in certain groups when the risk reductions for these groups were insignificant when analyzed using a random-effects model, including the cases with ulinastatin administration only prior to ERCP as well as high-risk patients, those receiving low-dosage ulinastatin and low-quality studies (data not shown), indicating the presence of substantial heterogeneities and inconsistencies among the studies included in these sub-groups.
Table II. Studies included in the meta-analysis on the incidence of PEP.

| Author, year       | Summary result                                                                 | Sample size (n) | Incidence of PEP (n/%)                  | Definition of PEP                                                                 |
|--------------------|--------------------------------------------------------------------------------|-----------------|----------------------------------------|-----------------------------------------------------------------------------------|
| Gong et al, 2004   | Ulinastatin significantly reduced PEP incidence vs. control (P<0.05).          | 68              | 5/7.35 vs. 9/14.52                     | PEP defined as serum amylase levels of >500U/l, accompanied by upper-middle abdominal pain lasting for >24 h. |
| Song et al, 2005   | Ulinastatin significantly reduced PEP incidence vs. control (P<0.05).          | 20              | 0/0 vs. 2/10.00                       | PEP defined as increased serum amylase levels lasting >24 h accompanied with upper abdominal pain, vomiting, nausea and the need for hospitalization. |
| Tsujino et al, 2005| Incidence of PEP was significantly lower in the ulinastatin group vs. control (P=0.041), and the severity of PEP was not significantly different between the 2 groups. | 204             | Total, 6/2.94; mild, 4/1.96; moderate, 2/0.98; severe, 0/0 | Acute PEP defined as abdominal pain persisting for at least 24 h after ERCP associated with a high serum amylase or lipase levels equivalent to at least 3 times the upper limit of normal 18 h after the procedure. Pancreatitis was graded according to a modification of the 1991 Consensus Guidelines: Mild, requiring fasting and treatment for 3 days or less; moderate, requiring fasting and treatment for 4-10 days; severe, requiring fasting and treatment for >10 days, intensive care, or surgical intervention. |
| Yoo et al, 2008    | No significant difference between the 2 groups (P=0.715).                       | 119; all high-risk patients<sup>a</sup> | Total, 8/6.72; mild, 7/5.88; moderate, 1/0.84; severe, 0/0 | Acute PEP was defined as the presence of abdominal pain typical for pancreatitis at 24 h post-ERCP with hyperamylasemia. Pancreatitis was graded according to the 1991 Consensus Guidelines as stated above. |
Table II. Continued.

| Author, year | Summary result | Sample size (n) | Incidence of PEP (n/%) | Definition of PEP | (Refs.) |
|--------------|----------------|----------------|-----------------------|------------------|--------|
|              |                | Ulinastatin    | Control               | Ulinastatin      | Control |          |
| Wang et al., 2010 | Incidence of PEP was significantly lower in the ulinastatin group vs. placebo (P<0.05). | 93 | 85 | 4/4.30 | 11/12.94 | PEP was diagnosed if serum amylase levels were at least 3 times the upper limit of normal at 3 and 24 h after ERCP accompanied by abdominal pain, fever or hospitalization for >3 days. (14) |
| Yoo et al., 2012 | Incidence of PEP in the ulinastatin group comparable to Control (P>0.05). | 229 | 658 | Total, 16/6.99 mild, 10/4.37; moderate to severe, 6/2.62 | Total, 48/7.29 mild, 33/5.02; moderate to severe, 15/2.28 | PEP was defined based on the 1991 Consensus Guidelines as stated above (24). The severity was graded as mild when hospitalization lasted 2-3 days, moderate when it lasted 4-10 days and severe when it lasted for >10 days or if any of the following conditions occurred: Hemorrhagic pancreatitis, pancreatic necrosis, pancreatic pseudocyst, or a need for percutaneous drainage or surgery. (4) |
| Xiong et al., 2013 | Incidence of PEP was significantly lower in the ulinastatin group vs. control (P<0.05). | 37 | 36 | 2/5.41 | 7/19.44 | PEP was diagnosed if serum amylase levels were at least 3 times the upper limit of normal after ERCP accompanied by abdominal pain and tenderness. (13) |
| Park et al., 2014 | Ulinastatin significantly reduced PEP incidence in overall population, (P<0.05) not in low-risk patients, but in high-risk patients (P<0.001). | Total, 53; low-risk, 11; high-risk, 42 | Total, 53; low-risk, 37; high-risk, 16 | Total, 1/1.89 (mild); 5/9.43; moderate, 2/3.77; severe, 0/0. All patients were high-risk. | Total, 7/13.21; mild, 5/9.43; moderate, 2/3.77; severe, 0/0. All patients were high-risk. | PEP was diagnosed when new-onset or increased abdominal pain lasted for 24 h, associated with an increase in serum amylase or lipase to at least 3 times the normal level ~24 h after the procedure. The severity was graded as mild when hospitalization lasted for 2-3 days, moderate with 4-10 days of hospitalization and severe for >10 days of hospitalization or when any of the following occurred: Hemorrhagic pancreatitis, pancreatic necrosis, pancreatic pseudocyst and the need for percutaneous drainage or surgery. (9) |
Table II. Continued.

| Author, year | Summary result | Sample size (n) | Incidence of PEP (n/%) | Definition of PEP | (Refs.) |
|--------------|----------------|----------------|------------------------|-------------------|--------|
|              |                | Ulinastatin | Control | Ulinastatin | Control |          |            |
| Wang et al, 2014 | Incidence of PEP was significantly lower in the ulinastatin group vs. placebo (P<0.05). | 160 | 120 | 7/4.38 | 13/10.83 | PEP was diagnosed if serum amylase levels were equivalent to at least 3 times the upper limit of normal 3 and 24 h after ERCP accompanied with abdominal pain, fever or hospitalization persisting for >3 days after ERCP. | (7) |
| Chen et al, 2015 | Ulinastatin significantly reduced the incidence of PEP vs. control (P<0.05). | 60 | 60 | 1/1.67 | 5/8.33 | PEP was diagnosed if hyperamylasemia accompanied by acute abdominal pain and vomiting were present, in addition to upper abdominal tenderness and rebound tenderness. Hyperamylasemia was defined as amylase levels higher than the upper limit of normal without abdominal pain or vomiting. | (16) |
| Chen et al, 2015 | Ulinastatin significantly reduced incidence of PEP vs. control (P<0.05). | 60 | 59 | 1/1.67 | 6/10.17 | PEP was diagnosed if abdominal pain, nausea, vomiting and upper abdominal tenderness and rebound tenderness lasting for at least 24 h, associated with an increase in serum amylase levels of at least 3 times the upper limit of normal were present. | (17) |

*High-risk patients were defined as those with any of the following conditions: i) Difficult cannulation (4 attempts or more); ii) pancreatic duct visualization with acinar filling; iii) a history of previous post-ERCP pancreatitis; iv) endoscopic pancreatic sphincterotomy or another pancreatic duct procedure, such as brush cytology or biopsy; v) pre-cut sphincterotomy; or vi) balloon dilatation of the bile duct. Patients were classified as high-risk if they had any of the following: Age <60 years, history of PEP, difficult cannulation, endoscopic pancreatic sphincterotomy or another pancreatic duct procedure, such as brush cytology or biopsy, precut sphincterotomy, or balloon dilatation of the bile duct. PEP, post-ERCP pancreatitis; ERCP, endoscopic retrograde cholangiopancreatography.*
Table III. Studies included in the meta-analysis on the incidence of PEHA.

| Author, year   | Summary result                                                                 | Sample size (n) | Incidence of PEHA (n/%) | Definition of PEHA                                                                 |
|----------------|-------------------------------------------------------------------------------|-----------------|-------------------------|-----------------------------------------------------------------------------------|
| Song et al, 2005 | Ulinastatin significantly decreased PEHA incidence vs. control (P<0.05).        | 20/20          | 2/10.00                 | PEHA was diagnosed with increased serum amylase levels without clinical symptoms and when the serum amylase levels returned to normal with 24 h. (11) |
| Tsujino et al, 2005 | Incidence of PEHA was significantly lower in the ulinastatin group vs. control (P=0.011). | 204/202        | 30/4.71                 | Hyperamylasemia was defined as amylase levels >3 times the upper time of normal at 4 or 18 h after ERCP. (6) |
| Yoo et al, 2008  | No significant difference between the 2 groups (P=0.510).                      | 119; 108        | 13/10.93                | Hyperamylasemia was defined as an elevation in serum amylase levels to >3 times the normal upper limit at 24 h post-ERCP. (8) |
| Wang et al, 2010 | No significant difference between the 2 groups (P>0.05).                        | 93; 85          | 36/38.71                | PEHA was defined as the elevation of serum amylase level to at least 3 times the upper limit of normal 3 and 24 h after ERCP without abdominal pain, fever or hospitalization for >3 days. (14) |
| Yoo et al, 2012  | No significant difference between the 2 groups (P>0.05).                        | 229; 658        | 25/10.92                | Hyperamylasemia was defined as elevation of serum amylase levels without abdominal pain and other abdominal symptoms. (13) |
| Xiong et al, 2013 | Incidence of PEHA was significantly lower in the ulinastatin group vs. control (P<0.05). | 37; 36          | 9/24.32                 | PEHA was defined as a 3-fold or greater increase in serum amylase level at 24 h after ERCP, without other symptoms. (9) |
| Park et al, 2014 | Ulinastatin significantly reduced PEHA incidence in overall population (P<0.05), not in low-risk patients, but in high-risk patients (P<0.001). | Total, 53; low-risk, 11; high-risk, 42 | Total, 53; low-risk, 37; high-risk, 16 | PEHA was defined as anylase level more the upper limit of normal level without abdominal pain or vomiting. (16) |
| Chen et al, 2015 | Ulinastatin significantly reduced the incidence of PEHA vs. control (P<0.05).    | 60/60           | 8/13.33                 | -                                                                                 |

*PEHA was defined as amylase level more the upper limit of normal level without abdominal pain or vomiting. (16)
Effect of prophylactic ulinastatin administration on preventing post-ERCP abdominal pain. As presented in Table IV, 7 studies including 582 patients receiving prophylactic ulinastatin and 590 patients receiving placebo or nothing as controls were included in the meta-analysis on the efficacy of prophylactic ulinastatin administration in preventing post-ERCP abdominal pain (6,8-10,15-17). The analysis revealed that overall, ulinastatin administration only marginally reduced the risk of post-ERCP abdominal pain vs. control (RR=0.67; 95% CI: 0.45-1.00; P=0.05; I²=67), however, pre-ERCP ulinastatin administration significantly reduced the risk of post-ERCP abdominal pain (RR=0.59; 95% CI: 0.42-0.83; P=0.002; I²=37), while ulinastatin administration commenced during or after ERCP did not (Fig. 4A). Furthermore, post-ERCP ulinastatin administration in addition to the pre-ERCP administration appeared to be necessary to significantly reduce the incidence of post-ERCP abdominal pain (RR=0.56; 95% CI: 0.35-0.89; P=0.01; I²=54; Fig. 4B).

In addition, patients with a low or average risk for PEP who received ulinastatin had a significantly reduced risk of post-ERCP abdominal pain vs. control (RR=0.59; 95% CI: 0.35-0.89; P=0.01; I²=20), but patients with a high risk for PEP did not (Fig. 4C). Furthermore, high-dose ulinastatin administration (150,000 U) led to a significantly reduced incidence of post-ERCP abdominal pain (RR=0.64; 95% CI: 0.44-0.93; P=0.02; I²=0), while low-dosage ulinastatin administration did not (50,000 or 100,000 U; Fig. 4D).

The present sub-group analysis based on the quality of the studies revealed that the quality had an impact on the results. In fact, although analysis including low-quality studies revealed a significantly reduced incidence of post-ERCP abdominal pain associated with ulinastatin vs. control, analysis including high or average-quality studies did not (Fig. 4E).

Using a fixed-effects model, the marginally significant reduction of the risk of post-ERCP abdominal pain associated with ulinastatin reached statistical significance (RR=0.71; 95% CI: 0.57-0.88; P=0.002; I²=67). Similar results were obtained for patients who received low-dosage ulinastatin and the sub-group only including low-quality studies, indicating heterogeneities and inconsistencies among these studies.

Heterogeneity and publication bias. Inter-study heterogeneities varied from none (I²=0) to very high (I²=96). Funnel plots for all of the analyses in the present study were slightly asymmetrical, wherein funnel plots for the analysis on the efficacy of prophylactic ulinastatin in preventing PEP were more symmetrical than those for the analysis on PEHA and post-ERCP abdominal pain (data not shown), indicating the possible presence of publication bias, with analysis on PEP having potentially less publication bias than those on PEHA and post-ERCP abdominal pain (data not shown).

Discussion

The present meta-analysis focused on the efficacy of prophylactic ulinastatin in preventing PEP, PEHA and post-ERCP abdominal pain. The analysis revealed that prophylactic ulinastatin significantly reduced the PEP risk vs. control; however, such significant risk reduction occurred only in patients with a low or average risk for PEP, with high-dosage ulinastatin...
Table IV. Studies included in the meta-analysis on the incidence of post-ERCP abdominal pain.

| Author, year   | Summary result                                                                 | Sample size (n) | Incidence of post-ERCP abdominal pain (n/%) | (Refs.) |
|---------------|---------------------------------------------------------------------------------|-----------------|--------------------------------------------|---------|
| $^{a}$Ohwada et al, 1997 | Ulinastatin significantly reduced the incidence of abdominal pain vs. control (P<0.05). | 22              | 6/27.27                                   | 20/43.48 | (10) |
| $^{b}$Tsujino et al, 2005 | No significant difference between the 2 groups (P=0.081). | 204             | 18/8.82                                   | 29/14.36 | (6)  |
| Yao et al, 2005 | Ulinastatin significantly reduced the incidence of abdominal pain vs. control (P<0.05). | 64              | 2/3.13                                    | 18/29.03 | (15) |
| Yoo et al, 2008 | No significant difference between the 2 groups (P=0.061).                        | 119; all high-risk patients $^{a}$ | 36/30.25                                   | 21/19.44 | (8)  |
| Park et al, 2014 | Ulinastatin significantly reduced the incidence of abdominal pain in overall population (P<0.05), not in low-risk patients, but in high-risk patients (P<0.001). | Total, 53; low-risk, 11; high-risk 42 $^{b}$ | Total, 16/30.19; low-risk patients, 2/3.77; high-risk patients, 14/26.42 | Total, 24/45.28; low-risk patients, 9/16.98; high-risk patients, 15/28.30 | (9)  |
| Chen et al, 2015 | No significant difference between the 2 groups (P>0.05).                          | 60              | 14/23.33                                   | 21/35.00 | (16) |
| Chen et al, 2015 | No statistical significance between the 2 groups (P>0.05).                        | 60              | 16/26.67                                   | 25/42.37 | (17) |

Notes:
- $^{a}$High-risk patients were defined as those with any of the following conditions: i) Difficult cannulation (4 attempts or more); ii) pancreatic duct visualization with acinar filling; iii) a history of previous post-ERCP pancreatitis; iv) endoscopic pancreatic sphincterotomy or another pancreatic duct procedure, such as brush cytology or biopsy; v) pre-cut sphincterotomy; or vi) balloon dilatation of the bile duct.
- $^{b}$Patients were classified as high-risk if they had any of the following: Age <60 years, history of post-ERCP pancreatitis, difficult cannulation, endoscopic pancreatic sphincterotomy or another pancreatic duct procedure, such as brush cytology or biopsy, precut sphincterotomy, or balloon dilatation of the bile duct. ERCP, endoscopic retrograde cholangiopancreatography.
Figure 2. Incidence of PEP in patients who received prophylactic ulinastatin vs. control. (A) Analysis with stratification by whether the ulinastatin administration began prior to, during or after ERCP. (B) Analysis for patients who began ulinastatin administration prior to ERCP, further stratified by whether ulinastatin was administered only prior to ERCP or also after ERCP. (C) Analysis with patients stratified into those with high risk for PEP and others.
Additional post-ERCP ulinastatin administration was unnecessary. Prophylactic ulinastatin administration also significantly reduced the risk of PEHA and post-ERCP abdominal pain in patients with a low or average risk for PEP, with high-dosage ulinastatin administration (150,000 or 200,000 U), and when ulinastatin administration began prior to ERCP with the help of additional post-ERCP ulinastatin administration.

The present results on the preventive effect of ulinastatin on PEP, PEHA and post-ERCP abdominal pain were consistent with those of the meta-analysis published by Chen et al (3) in 2010, which included 5 randomized controlled studies and found that ulinastatin prevented PEP and PEHA in average-risk patients when administered intravenously at a dose of ≥150,000 U. However, the results of the present study were inconsistent with the meta-analysis by Yuhara et al (1) from 2014, including (150,000 or 200,000 U) and when the ulinastatin administration began prior to or during ERCP. Additional post-ERCP ulinastatin administration was unnecessary. Prophylactic ulinastatin administration also significantly reduced the risk of PEHA and post-ERCP abdominal pain in patients with a low or average risk for PEP, with high-dosage ulinastatin administration (150,000 or 200,000 U), and when ulinastatin administration began prior to ERCP with the help of additional post-ERCP ulinastatin administration.

The present results on the preventive effect of ulinastatin on PEP, PEHA and post-ERCP abdominal pain were rather consistent with a couple of exceptions. Prophylactic ulinastatin administration also significantly reduced the risk of PEHA and post-ERCP abdominal pain in patients with a low or average risk for PEP, with high-dosage ulinastatin administration (150,000 or 200,000 U), and when ulinastatin administration began prior to ERCP with the help of additional post-ERCP ulinastatin administration.
Figure 3. Incidence of post-ERCP hyperamylasemia in patients who received prophylactic ulinastatin vs. control. (A) Analysis stratified by whether ulinastatin administration began prior to, during or after ERCP. (B) Analysis for patients who began ulinastatin administration prior to ERCP, further stratified by whether ulinastatin was administered only prior to or also after ERCP. (C) Analysis with patients stratified into those with high risk for post-ERCP hyperamylasemia and other patients.
6 randomized trials, two of which were high-quality studies with a Jadad score ≥3, which found that ulinastatin was not associated with a decreased PEP risk regardless of the quality of the studies. The present meta-analysis with 13 studies and a larger population size confirmed and further expanded on the findings by Chen et al (3). Since the present meta-analysis included several studies published after the study of Yuhara et al (1) and also several earlier-published Chinese studies not included by them, and since these studies were grouped into high- or average-quality (Jadad score, ≥2) and low-quality studies (Jadad score, <2). The squares and horizontal lines represent risk ratio and 95% CI for included studies, respectively. CI, confidence interval; M-H, Mantel-Haenszel; ERCP, endoscopic retrograde cholangiopancreatography; df, degrees of freedom.

Figure 3. Continued. (D) Analysis stratified by the dosage of ulinastatin administered: Low-dosage group (100,000 U) and high-dosage group (150,000 or 200,000 U). (E) Analysis stratified by the qualities of the included studies: High or average-quality studies (Jadad score, ≥2) and low-quality studies (Jadad score, <2). The squares and horizontal lines represent risk ratio and 95% CI for included studies, respectively. CI, confidence interval; M-H, Mantel-Haenszel; ERCP, endoscopic retrograde cholangiopancreatography; df, degrees of freedom.

PEP is the most common complication of ERCP and is potentially fatal. It starts with premature intra-acinar trypsin activation and subsequent activation of various proteolytic enzymes (6). Trypsin activates enzymes such as phospholipase A₂, kallikrein and elastase, and leads to injury of acinar cells, autodigestion and a systemic inflammatory response (6). Given that trypsin activation is the triggering event, ulinastatin, a trypsin inhibitor, makes a perfectly plausible candidate for PEP prevention. In addition, ulinastatin has the added advantage of being a strong inhibitor of a wide variety of pancreatic enzymes such as amylase, lipase and elastase, and also being able to modulate the immune response by suppressing pain was further analyzed. When there is no pain, there is no pancreatitis, and PEP only occurs in patients who complain of abdominal pain accompanied by hyperamylasemia (3,22). Therefore, similar to PEHA, the effect of prophylactic ulinastatin on post-ERCP abdominal pain should also be analyzed.
Figure 4. Incidence of post-ERCP abdominal pain in patients who received prophylactic ulinastatin vs control. (A) Analysis with stratification by whether the ulinastatin administration began prior to, during or after ERCP. (B) Analysis for patients who began ulinastatin administration prior to ERCP, further stratified by whether ulinastatin was administered only prior to ERCP or also after ERCP. (C) Analysis with patients stratified into those with high risk for post-ERCP abdominal pain and others.
pro-inflammatory mediators such as interleukin-6 and tumor necrosis factor-α, which are associated with the severity of pancreatitis (3,6,23). The present analysis supported the efficacy of ulinastatin in preventing PEP, although several conditions are required for this. First, the present analysis revealed that ulinastatin was only effective when its administration began prior to or during ERCP. Considering the fact that in pancreatitis, the acute-phase response to pancreatic injury starts immediately after pancreatic damage, the present result makes sense, since any attempt to prevent PEP must start sufficiently early so that any trypsin activation-induced downstream events are discontinued. Furthermore, the present analysis revealed that only high-dosage ulinastatin (150,000 or 200,000 U) was effective. Indeed, for a medication to work properly, a sufficient concentration in its target organ/tissue should be achieved, and in the scenario of the present study, a dosage of at least 150,000 U is needed. In addition, the present analysis revealed that prophylactic ulinastatin was only effective in patients with a low risk for PEP; however, only 2 studies including 161 patients receiving prophylactic ulinastatin and 124 patients taking a placebo were included for analysis of high-risk patients (8,9), and in one of the studies with a 72%-weight in the analysis, ulinastatin was administered after ERCP at a low dose (100,000 U) (8). As revealed by the present analysis, ulinastatin administered after ERCP or at a low dose was not effective; therefore, the question as to whether prophylactic ulinastatin may be effective in high-risk patients if administered timely and at a high dosage was not properly answered. The other study, by Park et al (9), found that ulinastatin significantly reduced the risk of PEP in high-risk patients; however, it was a small-scale study including only 42 high-risk patients receiving ulinastatin and 16 high-risk patients receiving placebo, and as such, its results are less than solid. The question as to whether ulinastatin is effective in these high-risk patients is important because after all, the overall frequency of PEP is relatively low (1-10%) and most PEP cases are mild (1,2); therefore, routine prophylactic ulinastatin administration for every patient undergoing ERCP...
does not appear to be cost-effective. However, if ulinastatin is effective in patients with a high risk for PEP, administration of ulinastatin only to high-risk patients may be more cost-effective. Although certain risk factors for PEP, including difficulty in cannulation, pre-cut sphincterotomy and pancreatic acinarization, were only identified during or after ERCP (8), other risk factors such as young age, female gender, suspected sphincter of Oddi dysfunction, previous PEP and therapeutic ERCP (9) were readily identifiable prior to ERCP. If prophylactic ulinastatin is effective in high-risk patients, pre-ERCP ulinastatin administration may reduce the risk of PEP for patients in which risk factors are identifiable prior to ERCP. The present analysis on timing of ulinastatin administration also revealed that starting ulinastatin administration during ERCP may significantly reduce the incidence of PEP; however, the present analysis only included 2 studies with 253 patients receiving ulinastatin and 205 patients receiving placebo (7,14); therefore, the validity of this result was less than definite and more studies on this topic are required. If starting ulinastatin administration during ERCP is not too late to prevent PEP, patients presenting with risk factors for PEP during ERCP, such as difficult cannulation and pre-cut sphincterotomy, may be administered high-dosage ulinastatin during ERCP to prevent PEP. In summary, it is worthwhile to perform additional studies on whether prophylactic ulinastatin is effective in preventing PEP in high-risk patients and also on whether starting ulinastatin administration during ERCP may still prevent PEP in order to improve the cost-effectiveness of prophylactic ulinastatin by administering ulinastatin only to high-risk patients.

The present analysis further revealed that post-ERCP ulinastatin administration in addition to pre-ERCP ulinastatin did not have any added benefit, and that pre-ERCP high-dosage ulinastatin was effective in preventing ERCP, so repeated ERCP administration is not necessary. However, since only 2 studies with 224 patients taking ulinastatin and 222 patients taking placebo were included (6,11), more studies on this topic are required to further confirm this result.

The present study also analyzed the preventive effect of ulinastatin on PEHA and post-ERCP abdominal pain, since they are possible symptoms of PEP. Asymptomatic PEHA occurs in 35-75% of patients undergoing ERCP, the degree of amylase elevation does not appear to correlate with the severity of PEP, and it is rather the timing of hyperamylasemia that has an important role in the early diagnosis of PEP (3,8). Normally, PEP develops between 3-7 h post-ERCP, and an increase in serum pancreatic enzymes such as amylase and lipase may be observed 2-4 h after ERCP and abdominal pain starts between 3-7 h post-ERCP (3,8,22). Therefore, studies on the effect of ulinastatin on preventing PEHA and abdominal pain should focus on reducing their incidence within a couple of h post-ERCP.

The present analysis was limited by several factors. First and foremost, inter-study heterogeneity was present, as reflected by the impact of using a fixed-effects model on certain results. These heterogeneities may be due to multiple factors. Definitions of PEP and PEHA as well as the time-point(s) for recording abdominal pain differed among the studies. Only unification of the definitions for PEP and PEHA and also the time-point(s) of recording abdominal pain will substantially decrease the heterogeneities. The 1991 Consensus Guidelines defined PEP as a newly developed or increased abdominal pain within 24 h after ERCP requiring analgesic agents, accompanied by the elevation of serum amylase and/or lipase levels at least 3 times the normal upper limit ~24 h after ERCP (24), and we recommend this definition of PEP. Another reason for such substantial heterogeneities may be the differences in study design. The present study made no adjustment for confounding effects of pancreatic stenting; recent improvements in endoscopic techniques contributed to a decreased risk of PEP in recent years, and as such, the beneficial effects associated with prophylactic ulinastatin in the later studies may be somewhat obscured.

The present study was further limited by the presence of low-quality studies, as reflected by the impact of excluding low-quality studies from the results on the effect of prophylactic ulinastatin in prevent PEHA and post-ERCP abdominal pain. However, sub-group analysis based on the quality of included studies was performed to determine the robustness and steadiness of the present results. Furthermore, possible publication bias was identified among the studies, as reflected by asymmetrical funnel plots.

At the same time, the present analysis had several strengths. First, all relevant studies not only published in English, but also in Chinese, were included, and considering the fact that a substantial portion of the studies on the effect of prophylactic ulinastatin on preventing PEP were performed in China, inclusion of studies published in Chinese improved the statistical power of the analysis. In addition, the quality of each study included was rigorously assessed and a sub-group analysis was performed based on the quality of the studies to test the validity and robustness of the present analyses. Furthermore, multiple outcome measures were used, and in addition to PEP and PEHA, the effect of ulinastatin on post-ERCP abdominal pain was also analyzed. Finally, multiple sub-group analyses were performed in order to further delineate a precise method of ulinastatin administration.

In conclusion, prophylactic ulinastatin administration significantly reduced the risk of PEP in patients with low or average risk for PEP when administered at a high dosage prior to or during ERCP. High-quality studies on high-risk patients and for ulinastatin administered during ERCP are warranted.

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