Pharmacokinetic Changes Secondary to Roux en Y Gastric Bypass

Christopher Giuliano1*, Sheila M. Wilhelm2 and Pramodini B. Kale-Pradhan1

1Eugene Applebaum College of Pharmacy and Health Science, Department of Pharmacy Practice, Wayne State University, St. John Hospital and Medical Center, Detroit, MI 48201, USA
2Eugene Applebaum College of Pharmacy and Health Science, Department of Pharmacy Practice, Wayne State University, Harper University Hospital, Detroit, MI 48201, USA

Abstract

Objective: To review the influence of Roux-en-Y gastric bypass (RYGB) on pharmacokinetic parameters of medications.

Data sources: PubMed was searched from inception to September 2012 to identify studies. Search terms included bariatric surgery, gastric bypass, Roux-en-Y, pharmacokinetic, and absorption. Studies included for this review were limited to English language studies published in full.

Data synthesis: Obesity is a major health care concern and is on the rise. This has led to an increasing number of bariatric surgeries. Such procedures may have profound effects on pharmacokinetic parameters of many medications depending on the extent of surgical changes that are made. Surgical procedures such as RYGB are most likely to affect medication absorption. Factors that may affect medication absorption in RYGB patients include changes in intestinal or gastric pH, surface area, intestinal metabolism and transport mechanisms. Published studies have been primarily conducted in RYGB patients and have shown varied effect on overall absorption of medications.

Conclusions: RYGB may have profound effects on medication absorption. Predicting absorption is difficult due to interplay of several factors including, changes in intestinal surface area, intestinal metabolism, efflux pumps, active transporters and gastrointestinal pH. Future studies are needed, particularly studies evaluating medications that have a low bioavailability and are commonly used in the bariatric surgery population.

Keywords: Roux en Y gastric bypass; Gastric; Intestine; Tmax; Cmax

Introduction

Obesity is a growing epidemic leading to worldwide public health concerns. In 1999–2000, 27.5% of men and 33.4% of women were obese [1]. The prevalence increased to 35.5% in men and 35.8% in women by 2009–2010 [1]. Bariatric surgery is an excellent option for patients with clinically severe obesity, defined as body mass index (BMI) of greater than 40 or BMI of >35 with serious comorbid conditions [2]. The number of bariatric surgical procedures has increased from approximately 13,000 in 1998 to over 121,000 in 2004 in the United States [3,4]. There are three types of procedures that are typically employed to address the issue of obesity including restrictive, malabsorptive and restrictive-malabsorptive. Each of these procedures may affect medication absorption differently based on the anatomical changes that are made during surgery.

Bariatric Procedures

Restrictive procedures significantly reduce the gastric capacity and limit oral intake thereby producing weight loss [5]. Common restrictive procedures include vertical banded gastroplasty, adjustable gastric banding, and sleeve gastrectomy. Vertical banded gastroplasty is performed by introducing a vertical partition in the stomach at the gastro esophageal junction to produce a gastric pouch [3]. The stoma between the gastric pouch and the remainder of the stomach is reinforced with a band to prevent dilation of the opening. This procedure has fallen out of favor due to development of persistent vomiting or gastro esophageal reflux and an inflammatory response leading to scarring. Adjustable gastric banding is a procedure that involves placing an adjustable silicone band 1-2 cm below the gastro esophageal junction which creates a 20-30 milliliter (mL) upper gastric pouch [6]. The constriction of the band may be adjusted using a saline injection in to a subcutaneous port. This procedure is easily reversible and is increasing in popularity. Sleeve gastrectomy is a procedure where the greater curvature of the stomach is removed [7]. This procedure can be used alone or combined with bilipancreatic diversion.

Malabsorptive procedures modify the length of the intestines, which decreases absorption of nutrients. Jejunileal or intestinal bypass is a surgical procedure that bypasses more than 90% of the small intestine without manipulating the stomach [3]. This is accomplished by forming a blind loop and connecting the proximal jejunum to the terminal ileum. This procedure has fallen out of favor because of related complications including metabolic issues, hepatic failure, nephrolithiasis and autoimmune complications.

Restrictive-malabsorptive procedures such as Roux-en-Y gastric bypass (RYGB) and bilipancreatic diversion combine both of these approaches. The RYGB is one of the more common gastric bypass procedures and has been the focus of trials effecting medication absorption [4]. It involves dividing the stomach to form a gastric pouch

*Corresponding author: Christopher Giuliano, Pharm. D, Assistant Professor (Clinical), Eugene Applebaum College of Pharmacy and Health Science, Department of Pharmacy Practice, Wayne State University, 259 Mack Avenue, Detroit, MI 48201, USA, Tel: (313) 343-4427; Fax: (313) 343-7632; E-mail: ek2397@wayne.edu

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with a capacity of 15 to 30 mL [3,6]. The jejunum is joined to the gastric pouch to form a gastro jejunanastomosis. This portion of the small intestine is the alimentary limb. The biliopancreatic limb includes the duodenum and a portion of the jejunum which is joined to the distal jejunum to form a jejunojejunostomy. The common channel is the portion of the small intestine distal to the jejunoojejunostomy. Biliary and pancreatic secretions deposit in the common channel allowing for nutrient digestion and absorption. The length of the common channel is a factor which determines the extent of absorption of nutrients (Figure 1). Biliopancreatic diversion is a procedure similar to the RYGB; however, it involves a partial gastrectomy rather than a division of the stomach. In addition, the biliopancreatic limb is diverted further distally than in RYGB. This produces a shorter common channel and a greater restriction of nutrient absorption. This review will examine the pharmacokinetic changes following restrictive malabsorptive procedures, with a specific focus on Roux-en-Y gastric bypass (RYGB).

**Pharmacokinetic changes after bariatric surgery**

Several factors that can affect medication pharmacokinetics may be altered after bariatric surgery. These changes are dependent on the type of surgical procedures. Restrictive procedures result in a greater restriction of nutrient absorption. This review will examine the pharmacokinetic changes following restrictive malabsorptive procedures, with a specific focus on Roux-en-Y gastric bypass (RYGB).

A major cause of decreased absorption in patients with bariatric surgery is reduced gastrointestinal surface area. The portion of the small intestine bypassed determines the effect on absorptive capacity. Moving from the duodenum to the ileum, the absorptive capacity of the small intestines decreases where as the transit time increases [2]. Although the proximal small intestine has the highest absorptive capacity, it has the shortest transit time. This may result in less overall absorption proximally compared to distal small intestines. Procedures that bypass the duodenum and proximal jejunum such as RYGB may have less of an impact on absorption compared to procedures that bypass the jejunum and the majority of the ileum such as the jejunoileal bypass. Also, medication absorption may be dependent on passive diffusion, active transport, paracellular transport, intracellular metabolism, efflux, and enterohepatic recirculation.

Passive diffusion relies on medication lipophilicity and concentration of medication in solution within the intestinal lumen relative to systemic concentration. Passive diffusion may occur anywhere along the small intestine, and to a lesser extent in the colon. The partition coefficient or Log P compares a medication’s preference to be in a hydrophobic environment such as octanol to a hydrophilic environment such as water [13]. Therefore, the higher the Log P of a medication, the more lipophilic and less soluble it is. Optimal log P values to facilitate both dissolution and passive diffusion are approximately 1-2 [14].

Unlike passive diffusion, active transport pumps facilitate movement of medication across intestinal cell membranes. Organic anion transporting polypeptides (OATP2B1 and OATP1A2), monocarboxylic acid transporter 1 (MCT1), and oligopeptide transporter (PEPT-1) are active transport pumps found primarily in the duodenum and jejunum [13]. Plasma membrane monoamine transporter (PMAT) is a transport pump with unknown distribution in the small intestines [15]. Medications with small molecular weight may undergo paracellular transport via tight junctions, although this process is saturable.

Intracellular enzymes such as CYP3A4, CYP2C9, CYP2C19, CYP2D6, phase 1 and 2 enzymes, uridine diphosphate, glucuronosyltransferases (UGT), phenylsulphotransferases (PST), and glutathione S-transferases (GST), are prevalent in the proximal small intestine and play a role in the metabolism of medications as they are absorbed [16]. Of the intestinal CYP enzymes, CYP3A is the most prevalent and accounts for more than 80% of intestinal CYP activity [17]. Some medications require enzymatic effects to be converted to their systemically active metabolite while others become inactivated by these enzymes. Medications may also be affected by the efflux pumps such as P-glycoprotein (PgP), breast cancer resistance protein (BCRP) and multidrug resistance associated protein 2 (MRP2) at the apical membrane of intestinal epithelial cells. These pumps transport intracellular molecules back into the intestinal lumen thereby reducing systemic absorption. PgP concentration increases as substances move from the duodenum to the ileum, which may decrease medication absorption following RYGB [18].

Medications that undergo enterohepatic recirculation are repeatedly exposed to the gastrointestinal tract [19]. In a non-surgical patient, a medication enters the gastrointestinal tract and is systemically absorbed. A portion of the systemically absorbed medication enters the

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**Figure 1: Roux-en-Y Gastric Bypass and factors that may influence medication absorption.**
liver and is partitioned into the bile where it re-enters the duodenum for repeated absorption [20]. Post-RYGB, the medication will be exposed to reduced surface area resulting in decreased initial and repeated absorption. Serum bile acid concentrations may be increased in RYGB patients [21]. Certain medications such as lovastatin and pravastatin have shown increased absorption with increasing concentrations of bile acids [22,23]. However, bile acids are not in high concentrations in the alimentary limb so the overall effect is unclear.

Overall the above mechanisms work individually or in combination to affect medication absorption. The interplay of these mechanisms makes it difficult to predict the effect of RYGB on the absorption of medications. This review will examine the available literature regarding RYGB and pharmacokinetic changes and in particular medication absorption.

Literature review

A search of PubMed from inception to September 2012 was conducted using the following key terms: bariatric surgery, gastric bypass, Roux-en-Y, pharmacokinetic, and absorption. Studies included for this review were limited to English language studies published in full. Bibliographies of recent relevant articles were hand searched to identify any additional studies. We found a total of 5 pharmacokinetic studies evaluating 7 medications in our literature search. These trials focused on both commonly used medications (sertraline, azithromycin, atorvastatin, metformin, tacrolimus, sirolimus, and mycophenolate mofetil). Specific properties that may have an effect on absorption of these medications are presented in Table 1.

In a prospective, case controlled study of sertraline pharmacokinetics, five subjects who had undergone RYG 9-15 months prior to enrollment were matched with five non-RYG subjects for gender, age and BMI [24]. Subjects were excluded from the study if they were found to be ultra-rapid or poor metabolizers for the CYP2D6 or 2C19 metabolic enzymes or were receiving any medication that is known to interact with sertraline. The primary endpoint was sertraline area under the curve (AUC) from 0-10.5 hours after oral ingestion of a single dose of sertraline 100 mg. Secondary endpoints were time to peak plasma level (Tmax) and maximum plasma concentration (Cmax). The RYGB and non-surgical groups were well matched for age (mean age 45.4 and 44.6 years, respectively) and BMI (29.9 and 30.6 kg/m², respectively). The mean AUC from 0-10.5 hours was significantly lower in the RYGB group compared to the non-surgical group (124.4 ng-hr/mL vs. 314.8 ng-hr/mL, respectively, p=0.043). The Cmax was also significantly less in the RYGB group compared to the non-surgical group (19 ng/mL vs. 48.7 ng/mL, respectively, p=0.043) but the Tmax was not significantly different (3.9 hr vs. 3.4 hr, respectively, p=0.357). Overall, this study was well conducted, although a longer evaluation of the AUC may have been more appropriate.

These changes may be partially predicted when assessing pharmacokinetic properties of sertraline. First, sertraline is a weak base with a pKa of 8.5 and exists primarily in the ionized state in the small intestine. RYGB should not affect the degree of ionization of sertraline. Due to its ionized state, sertraline likely requires active transport for absorption, although no active transporters have been identified. Secondly, in a simulated environment, sertraline undergoes significantly greater dissolution in a normal environment versus a RYG environment (16% dissolved vs. 10% dissolved, p<0.04) [25]. As the Log P value increases, solubility of medications decreases. A Log P of 4.8 in this situation reinforces that medication absorption is limited by its solubility [26]. Lastly, sertraline undergoes CYP2C19 and to a smaller degree of CYPD6 metabolism in the gut and also PgP excretion [27]. CYP2C19 and CYPD6 concentrations decrease as sertraline moves from the stomach to terminal ileum while PgP increases [18,28]. In an RYGB patient, this could lead to increased absorption as some intestinal metabolism is bypassed; however, the length of intestine likely plays a more important role as gastrointestinal CYP2C19 and especially CYPD6 have limited impact on sertraline metabolism.

A prospective pharmacokinetic study of azithromycin was completed in 14 female patients who had undergone RYG compared to 14 female non-surgical patients who were matched for BMI [29]. Enrolled patients were administered azithromycin 500 mg orally followed by plasma azithromycin sampling. The primary endpoint was the azithromycin AUC from 0-24 hours. Secondary outcomes were Cmax and Tmax. In the RYGB and non-surgical groups, mean age (44.1 years vs. 44.5 years, respectively, p=0.93) and BMI (36.8 vs. 35.9 kg/m², respectively, p=0.98) were similar between the two groups. AUC was significantly lower in subjects who had undergone RYG compared to non-surgical subjects (1.41 vs. 2.07 mg-hr/L, respectively, p=0.008).

<table>
<thead>
<tr>
<th>Medication</th>
<th>pKa</th>
<th>Lipophilicity (log P)</th>
<th>Intestinal metabolism</th>
<th>Efflux pump</th>
<th>Active transport pumps</th>
<th>Enterohepatic recirculation</th>
<th>Tmax (hr)</th>
<th>Bioavailability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sertraline</td>
<td>8.5</td>
<td>4.8</td>
<td>CYP2C19</td>
<td>PgP</td>
<td>No</td>
<td>No</td>
<td>4-6</td>
<td>44%*</td>
</tr>
<tr>
<td>Azithromycin</td>
<td>8.74</td>
<td>4</td>
<td>No</td>
<td>PgP</td>
<td>No</td>
<td>No</td>
<td>2-3</td>
<td>37%</td>
</tr>
<tr>
<td>Atorvastatin</td>
<td>4.46</td>
<td>1.53-acid</td>
<td>4.2-lactone*</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>5%</td>
</tr>
<tr>
<td>Metformin</td>
<td>11.5</td>
<td>-1.43</td>
<td>No</td>
<td>CYP3A4</td>
<td>No</td>
<td>No</td>
<td>3c</td>
<td>55%</td>
</tr>
<tr>
<td>Tacrolimus</td>
<td>14.07</td>
<td>2.7</td>
<td>CYP3A4</td>
<td>No</td>
<td>No</td>
<td>0.5-1</td>
<td>0.5-3</td>
<td>25% (5-93%)</td>
</tr>
<tr>
<td>Sirolimus</td>
<td>13.37</td>
<td>8</td>
<td>CYP3A4</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>1-2</td>
<td>15%</td>
</tr>
<tr>
<td>Mycophenolate Mofetil</td>
<td>15.67</td>
<td>3.2</td>
<td>UGT</td>
<td>MRP-2</td>
<td>PgP</td>
<td>Yes</td>
<td>80.7-94%</td>
<td></td>
</tr>
</tbody>
</table>

Table 1: Pharmacokinetic properties of medications in Roux-en-Y gastric bypass surgery.
Cmax was similar between the RYGB and nonsurgical groups (0.26 vs. 0.363 mg/dL, respectively, p=0.08) as was the Tmax (2.14 vs. 2.36 hours, respectively, p=0.75).

These changes may be partially predicted by the pharmacokinetic properties of azithromycin. Azithromycin is a weak base with a similar pKa and log P to sertraline, therefore its solubility will be a limiting factor in its absorption [26]. Azithromycin will be more likely to exist in its ionized state in the less acidic gastrointestinal tract following RYGB given its pKa. This hinders its passive diffusion, and an active transport mechanism for azithromycin has not been identified. Azithromycin is a substrate for the PgP efflux pumps, which are most prevalent in the distal small intestine and into the colon [30]. Following RYGB, azithromycin will be exposed to areas of high intestinal PgP activity which, when combined with reduced surface area, results in an overall decrease in systemic bioavailability.

The effects of RYGB on the pharmacokinetic parameters of atorvastatin were investigated in 12 patients (66% female, mean age 52 years) who were going to undergo RYGB [31]. Subjects who were scheduled for RYGB were eligible for the study if they were being treated with atorvastatin 20-80 mg daily (mean dose 37 mg daily). Atorvastatin kinetic evaluations were completed the day prior to surgery and again three to six weeks following surgery. The mean AUC from 0-8 hours did not significantly differ from prior to post RYGB (75 vs. 50 ng·hr/mL, respectively, p=0.99). Cmax and Tmax were also similar at both time points (Cmax 28 ng/mL vs. 13 ng/mL, respectively, p=0.83; Tmax 1.6 hr vs. 1.8 hr, respectively, p=0.39). The authors assessed the results for the patients with the highest and lowest systemic availability prior to surgery. The three patients with the highest AUC prior to surgery experienced a significant reduction post RYGB (median ratio of AUC 0.4, p=0.01) whereas the nine patients with the least AUC prior to surgery experienced a significant increase in AUC following RYGB (median ratio of AUC 1.2, p=0.03). This study is limited by the time frame used to assess pharmacokinetic changes. Patients had very recently undergone RYGB, and pharmacokinetic changes may vary as more time passes.

Atorvastatin converts to either acidic or lactone forms which make it both highly soluble and readily absorbed across the gastrointestinal tract [32]. Despite both high solubility and absorption, atorvastatin displays a bioavailability of 14% because of first pass metabolism in the liver and intestine as well as excretion by a variety of efflux pumps. Although atorvastatin undergoes active transport, this is unlikely to play a large role in its absorption [32]. Changes seen in this study may be explained by inter-individual variability in intestinal CYP3A4 metabolism. Intestinal CYP3A4 levels can vary up to a 20 fold [17]. In an individual with intrinsically low intestinal CYP3A4 activity, the loss of CYP3A4 exposure after surgery may not have a large effect on absorption, and rather, surface area may have a significant impact on medication absorption. Conversely, an individual with intrinsically high intestinal CYP3A4 activity may have an increase in absorption of atorvastatin because less medication will be metabolized in the small intestines.

A prospective, nonblinded trial assessed the pharmacokinetic parameters of metformin in 16 non-diabetic RYGB patients and 16 non-surgical patients matched for age (mean age 40.2 years) and BMI (mean BMI 39.2 kg/m²) [11]. The primary endpoint was the AUC of metformin from time zero extrapolated to infinity (0–∞). The secondary outcomes were Cmax, Tmax, bioavailability estimated from metformin excretion in the urine over 24 hours, and AUC of metformin from 0–24 hours. Nearly the entire bioavailable dose of metformin is eliminated within 24 hours; therefore, unlike most medications, 24 hour urinary excretion can be used to reliably estimate bioavailability without a required intravenous dose. The AUC 0–∞ was similar between the two groups, 13.7 mcg·hr/ml for RYGB subjects compared to 11.4 mcg·hr/ml for non-surgical patients (p=0.2). The Cmax was 2 mcg/ml versus 1.8 mcg/ml for RYGB and control, respectively (p=0.32). Tmax was 3 hours in both groups (p=0.89). The estimated bioavailability was significantly higher in the RYGB group compared to control (41.8% vs. 27.8%, p=0.007). The AUC from 0–24 hours was 13.4 and 11.1 mcg·hr/ml, respectively (p=0.2).

Pharmacokinetic properties of metformin partially explain the changes in AUC observed in the study. Metformin is a weak base with a pKa of 11.5, which exists primarily in a cationic state in the gastrointestinal tract. Its log P value of -1.43 also indicates that metformin has poor lipophilicity and will be unable to easily cross the gastrointestinal membrane. Both of these factors indicate that metformin likely undergoes active transport to be absorbed. PMAT and organic cation transporters 1 and 3 play a role in the absorption of metformin in the intestines with PMAT playing the most substantial role [15]. PMAT is a saturable transporter in the small intestine and may be a rate limiting step in the absorption of metformin. In RYGB patients gastric residence time will be increased allowing for slower release of metformin into the small intestine. This will result in less saturation of PMAT, which may lead to an increase in absorption.

In a pilot study of six patients who had undergone RYGB between 2 months and 7.41 years prior to the trial, pharmacokinetics of mycophenolatemofetil (MMF), sirolimus and tacrolimus were assessed [33]. Four of the six included patients had end stage renal disease requiring dialysis and were awaiting transplant. These patients received 24 hours of immunosuppressive therapy with MMF, sirolimus and tacrolimus. Two of the six patients had undergone renal transplant and were maintained on their immunosuppressive regimens. The kinetic parameters after the second dose of MMF in the three end stage renal disease patients with reported data included a mean AUC from 0-12 hours of 907.4 mcg·hr/L, a mean Cmax of 67.2 mcg/L, and a mean Tmax of 6.7 hours. For the two post-transplant patients, mean AUC from 0-12 hours was 903.9 mcg·hr/mL, mean Cmax was 69.8 mcg/mL, and mean Tmax was 2.5 hours. The kinetic parameters after a single dose of sirolimus in the four end stage renal disease patients were a mean AUC from 0-24 hours of 145.7 mcg·hr/L, AUC from 0–∞ of 181.1 mcg·hr/mL, a mean Cmax of 18.2 mcg/L, and a mean Tmax of 1.8 hours. Sirolimus was not assessed in the post-transplant patients. The kinetic parameters after the second dose of tacrolimus in the three end stage renal disease patients with reported data included a mean AUC from 0-12 hours of 58.8 mcg·hr/L, a mean Cmax of 12.3 mcg/L, and a mean Tmax of 3.3 hours. For the one post-transplant patient receiving tacrolimus, AUC from 12-24 hours was 63.8 mcg·hr/L, Cmax was 16 mcg/L, and mean Tmax was 1 hour. The study design did not include a comparator group of patients who had not undergone RYGB, which makes it difficult to draw conclusions about the potential changes in pharmacokinetic parameters due to RYGB. As well only six patients were enrolled in the study and not all were evaluable.

Pharmacokinetic changes in this study are difficult to predict due to the above mentioned limitations. Both tacrolimus and sirolimus have a wide variability in non-surgical patients and this may be more pronounced in RYGB patients [34,35]. Both medications have similar pKa's and will exist primarily in the ionized state, although the higher log P value for sirolimus may indicate less solubility than tacrolimus. Both medications are substrates for PgP efflux pumps and CYP3A4 intestinal metabolism.
metabolism. Similar to atorvastatin, absorption may be variable due to amounts of intestinal CYP3A4. Overall, the pharmacokinetic profiles suggest that there may be no change or a reduction in systemic bioavailability of sirolimus and tacrolimus.

MMF is an ester prodrug, which is quickly hydrolyzed in the gastrointestinal tract, blood, tissues, and liver to its active metabolitomecytophenolic acid (MPA) [36]. MPA undergoes glucuronidation by UGT in the small intestines, liver, and kidney and is also excreted by MRP-2 and PgP. UGT content is highest in the jejunum and ileum, which would not be affected by RYGB [37]. MPA also undergoes enterohepatic recirculation, exposing the shortened length of the intestine to the medication repeatedly. This aids in absorption, but to a lesser degree than it would in a patient with an intact gastrointestinal tract and greater available intestinal surface area. All of these changes would lead to an expected decrease in absorption for MMF following RYGB, which was observed in this study compared to historical controls.

Conclusion

Obesity is an epidemic that is receiving increasing attention as associated health issues are becoming more prevalent and we seek ways to effectively address these issues. Bariatric surgery and specifically RYGB is being used more commonly to treat obesity. Making significant changes to the gastrointestinal tract has ramifications on medication absorption. There are many factors that affect medication absorption including medication-specific characteristics such as pKa and Log P. In addition, patient specific factors such as expression of intestinal metabolic enzymes can affect medication absorption. Surgical changes to the gastrointestinal tract affect medication absorption through pH alteration, increased or decreased gastric residence time, surface area reduction, and altered exposure to metabolic enzymes and efflux pumps. Some of these changes may have a predictable effect on the absorption of a specific medication given its pharmacokinetic parameters; however it is difficult to accurately predict absorption following RYGDB due to the considerable interplay of these factors. In order to most effectively and safely treat patients that undergo RYGB, studies are needed to assess changes in absorption of medications in this population, particularly for medications with low bioavailability that are commonly used in the bariatric surgery population.

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