Stability of octreotide acetate decreases in a sodium bisulfate concentration-dependent manner: compatibility study with morphine and metoclopramide injections

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ABSTRACT

Purpose Sodium bisulfate is known to affect the stability of octreotide. However, the critical concentration of sodium bisulfate is not known. Therefore, we assessed the critical concentration of sodium bisulfate needed to preserve the stability of octreotide using actual drugs containing sodium bisulfate.

Methods Although morphine and metoclopramide preparations are considered to be compatible with octreotide, some of their products are known to contain sodium bisulfate. Thus, octreotide was mixed with preparations of sodium bisulfate solutions at serial concentrations and morphine and metoclopramide preparations containing sodium bisulfate, and octreotide stability was then evaluated using high performance liquid chromatography.

Results Octreotide concentrations decreased significantly at a sodium bisulfate concentration of 0.1 mg/mL or higher after 10 days when octreotide was mixed with sodium bisulfate solutions at various concentrations. A significant decrease in octreotide concentrations also occurred when it was mixed with morphine and metoclopramide preparations containing sodium bisulfate and stored for 10 days; however, slight decreases were observed in the mixture with both preparations and were within the clinically acceptable range for morphine preparations.

Conclusions These results indicate that the residual rate of octreotide decreases with time in a sodium bisulfate concentration-dependent manner when octreotide was mixed with morphine or metoclopramide. However, this incompatibility may be clinically acceptable when the final sodium bisulfate concentration is lower than 0.1 mg/mL and the mixed solution is used within 7 days.

INTRODUCTION

Carcinomatous peritonitis-induced gastrointestinal obstruction causes various digestive symptoms, such as nausea, vomiting and anorexia, which reduce the quality of life of patients. Octreotide (OCT) acetate is a somatostatin analogue that promotes the absorption of digestive juices and inhibits their secretion, and is also a key drug for treating digestive symptoms.1,2 OCT is a cyclic synthetic octapeptide in which the retention of the S–S bond and cyclic structure is necessary for it to exhibit its bioactivity (figure 1). Therefore, it contains an amide bond that connects intramolecular amino acids and is readily hydrolysed. Hydrolysis was shown to be promoted in alkaline solutions, with an increase in the inactivation speed being reported in the presence of sodium bisulfate (SBS) as an additive.3-8 We previously investigated incompatibilities with dexamethasone (DEX) injections, which are sold as an ethical drug, and demonstrated that OCT concentrations were maintained for 10 days in a mixture with a DEX injection that did not contain SBS but were significantly decreased in a mixture with a SBS-containing DEX injection.9 However, the concentrations of SBS that have a clinical influence have not yet been determined, and the stability of these drugs, other than DEX, remains unclear.

Morphine (MOR) 1% and 4% are sold in the form of injections in Japan; however, SBS is used as a stabiliser in the 4% preparation only because the concentration of MOR hydrochloride is close to its solubility, 0.178 mM (about 5.7% (W/V)).10 Although metoclopramide (MET) injections generally contain SBS or other stabilisers (such as sodium citrate) as an additive to stabilise the pH, this depends on the manufacturers. Previous studies have demonstrated that MOR and MET are compatible with OCT because no changes were observed in appearance, such as precipitation; however, the presence or absence of SBS as well as the hydrolysis of OCT were not described.11-13 Accordingly, the presence or absence of SBS and its concentration after mixing with OCT are considered to be very important factors in clinical studies evaluating the effects of OCT; they have been shown to directly influence the effects of OCT. In addition, the concentration of SBS that influences the stability of OCT has not been clarified, which presents difficulties when evaluating compatibilities with drugs containing SBS. Although MOR and MET are frequently used in combination with OCT for the palliative care of patients with end-stage cancer,12 to the best of our knowledge, no study has investigated compatibility with a focus on the presence or absence of SBS. In the case of the patients who self-administer analgesic treatment by using a patient-controlled analgesia pump (PCA pump: balloon type or portable mechanical type), the elucidation of the drug combination stability in the PCA pump would minimise the number of venipunctures and the usage of consumables, such as medical cassettes. Therefore, it
is advantageous both economically and for the patient’s quality of life.4–19

When aggressive treatment is not an option, OCT is an important drug that can be used to maintain the quality of life of patients until the end stage.1,19 However, large variations have been reported in the efficacy of OCT (44%–100%).18 The time after the occurrence of intestinal obstruction, obstructed region, infusion volume and presence or absence of a nasogastric tube and concomitant antimetic (such as MET and DEX) have already been proposed as factors that influence the effects of OCT;18 however, we hypothesised that this variation may be attributed to the SBS concentration. If this can be demonstrated, OCT can be used more effectively.

In the present study, we examined reproducibility, with a focus on the SBS concentration-dependent OCT stability of drugs made from MOR and MET, using high performance liquid chromatography (HPLC).

METHODS

Materials

OCT was obtained from Sigma-Aldrich Japan (Tokyo, Japan). SBS was obtained from Wako (Osaka, Japan). Sandostatin for subcutaneous injection (100 μg/1 mL ampoules) was purchased from Novartis Pharma (Tokyo, Japan). To obtain MOR products with different SBS concentrations, MOR hydrochloride injection 10 mg/1 mL ampoules (SBS free; 1% MOR–SBS 0) and MOR hydrochloride injection 200 mg/5 mL ampoules (SBS 0.1 mg/mL; 4% MOR–SBS 0.1) were purchased from Takeda Pharmaceutical (Tokyo, Japan). For MET products with different SBS concentrations, Elieten injection 10 mg/2 mL ampoules (SBS free; MET–SBS 0) and Primperan injection 10 mg/2 mL ampoules (SBS 1.09 mg/mL as sodium pyrosulphite 1 mg/mL; MET–SBS 1.0) were purchased from Takata Seiyaku (Tokyo, Japan) and Astellas Pharma (Tokyo, Japan), respectively. Sodium pyrosulphite is a dehydrating condensation compound of binary molecular SBS, the actions of which are similar to SBS in aqueous solution. All other chemicals were commercial products of analytical reagent grade.

Preparation of samples

Dependence on SBS concentration

SBS was dissolved and diluted with saline. Sample solutions were collected from an OCT ampoule and mixed with SBS. The mixture was diluted with saline to adjust the final SBS concentration to 0, 0.01, 0.03, 0.05, 0.1, 0.3, and 0.5 mg/mL and the final OCT concentration to 41.7 μg/mL, and 1 mL of each mixture was prepared. Each sample was added to a 1.5 mL microtube (Promega, Tokyo, Japan) and stored at room temperature with protection from light. OCT concentrations were then measured in samples containing SBS at various concentrations after 0, 3, 7 and 10 days using HPLC.

Compatibility study with MOR preparations

Based on the actual regimen, the experiment was performed on the assumption of the 7-day administration of daily doses of OCT and MOR at 300 μg and 20 mg, respectively.16 The necessary volumes were sampled from OCT and MOR ampoules and diluted with saline to a total volume of 50.4 mL. The final OCT and MOR concentrations were 41.7 μg/mL and 4.17 mg/mL, respectively. Two mixtures with 1% MOR–SBS 0 and 4% MOR–SBS 0.1 were prepared. The final SBS concentrations were 0 and 0.01 mg/mL, respectively. A total of 1 mL was transferred from each mixture to a 1.5 mL microtube and stored at room temperature with protection from light. The OCT concentration in each sample was then measured after 3, 7 and 10 days using HPLC.

Compatibility study with MET preparations

Based on the actual regimen, the experiment was performed on the assumption of the 7-day administration of daily doses of OCT and MET at 300 μg and 20 mg, respectively.16 As the concentration of MET was lower than that of MOR, the final volume, after mixing, may have exceeded the capacity of the smallest cassette reservoir (50 mL) of the initially prepared device (CADD-Legacy PCA; Smiths Medical Japan, Tokyo, Japan); therefore, it was necessary to consider the possibility of using a larger cassette (100 mL) than that used for MOR. The necessary volumes were sampled from OCT and MET ampoules and diluted with saline to a total volume of 101.0 mL. The final OCT and MET concentrations were 20.8 μg/mL and 1.39 mg/mL, respectively. Two mixtures with MET–SBS 0 and MET–SBS 1.0 were prepared. The final SBS concentrations were 0 and 0.30 mg/mL, respectively. A total of 1 mL was transferred from each mixture to a 1.5 mL microtube and stored at room temperature with protection from light. The OCT concentration in each sample was then measured after 3, 7 and 10 days using HPLC.

HPLC analysis

The HPLC device (LC-2010CHT; Shimadzu Corporation, Kyoto, Japan) and analysis software (LC solution; Shimadzu Corporation, Kyoto, Japan) were used in this analysis as described previously.19 The injection volume was 10 μL and detection wavelength was 210 nm (UV–VIS detector). Measurements were performed at room temperature using the reverse-phase column, COSMOSIL5C18-MS-II (column diameter: 4.6, column length: 150 mm) (Nacalai Tesque, Kyoto, Japan). Mobile phase A (water:acetonitrile:1 M tetramethylammonium hydroxide solution=440:50:10), mobile phase B (water:acetonitrile:1 M tetramethylammonium hydroxide solution=190:300:10) and mobile phase C (water:acetonitrile=50:50) were prepared, and the pH was set to 4.5 using two types of phosphate solution (0.17 M sodium dihydrogenphosphate and 0.17 M disodium hydrogenphosphate).

Using the linear gradient elution method, the mobile phase ratio was changed from A:B=73:27 to 55:45 at the initiation of analysis to 12 min, and the solution was allowed to flow at 1 mL/min to set C at 100% from 12 to 17 min. A calibration curve in the range of 0–50 μg/mL OCT was prepared from the peak height, and the OCT concentration was calculated from the peak height at the OCT retention time-point in each sample. We prepared a sample at every measurement point. After storing the samples, we assayed each sample three times. For
example, in the ‘dependence on SBS concentrations’ experiment, we prepared SBS at seven different concentration levels and assayed them on days 0, 3, 7 and 10. Therefore, we prepared samples of 28 measurement points (7 different levels × 4 days = 28 samples) and assayed them three times (28 samples × 3 times = 84 points measured).

Statistical analysis
Statistical comparisons were made using a one-way ANOVA (analysis of variance) followed by Tukey’s post-hoc test. Values of p<0.05 were considered to be significant. The clinical stability of OCT was defined as maintaining at least 90% of the initial OCT concentration.

RESULTS

Dependence on SBS concentrations
When the SBS concentration was lower than 0.1 mg/mL, no significant decrease was noted in the OCT concentration for 10 days after mixing with SBS at any concentration, and more than 90% of the initial OCT concentration was maintained (figure 2). However, the OCT concentration significantly decreased to less than 90% of the initial concentration on days 10, 3 and 0 after mixing when the concentrations of SBS were 0.1, 0.3 and 0.5 mg/mL, respectively.

Compatibility study with MOR preparations
The OCT concentration decreased significantly after storage for 10 days when OCT was mixed with 4% MOR containing SBS; however, more than 90% was maintained (figure 3). No significant decrease was noted in any combination in the mixture with 1% MOR.

Compatibility study with MET preparations
The OCT concentration decreased significantly after storage for 3 days and thereafter when OCT was mixed with MET–SBS 1.0. The OCT level was lower than 90% of the initial concentration at all time-points (figure 4). By contrast, no significant decrease was noted in any combination in the mixture with MET–SBS 0.

DISCUSSION

This is the first study to clarify the dependence of OCT stability on the SBS concentration, and the following results are of clinical importance.

SBS may not influence OCT stability when its final concentration is 0.05 mg/mL or lower; it may clinically influence stability and have a negative impact at a final concentration of 0.3 mg/mL or higher. A significant decrease was observed in the concentration of OCT on day 0, and this has been attributed to the progression of hydrolysis at the time of the measurement due to sample preparation time. Further decreases may occur in OCT concentration depending on other factors, such as the pH, temperature and exposure to light; therefore, mixing should be avoided under these conditions. Judgements regarding expiry dates (from immediately after preparation to the completion of its use) are varied when the final SBS concentration is 0.1 mg/mL. Although OCT concentrations decreased significantly on days 0 and 3, more than 90% of the initial concentration (theoretical value) was retained. Thus, these decreases may have no clinical influence when the preparation is used within 3 days. However, the stability of OCT may be influenced by SBS for its administration over a period that ranges from several days to a week.

Incompatibility with the SBS-containing MOR preparation was within a clinically acceptable range. SBS is only added to the 4% preparation to maintain the stability of MOR because the concentration is close to its solubility. Mixing with the 1% preparation was previously shown to be possible, but it was unclear whether it was possible with the 4% preparation because the presence of SBS had not been considered in previous compatibility studies. Our study demonstrated that, although the OCT concentration decreased significantly on day 10, more than 90% of the initial concentration (calculated value) was retained, indicating that this decrease had no clinical impact.
with OCT: the stability of OCT can be ensured by either selecting a formulation containing no SBS or verifying the SBS concentration; the limitation to patient activities are preventable by avoiding OCT loss and administration through another route. Staggered administration is unnecessary with mixing, thus possibly decreasing the frequency of venipuncture. Additionally, more reliable clinical studies may be needed to investigate the effects of OCT.

CONCLUSION

The stability of OCT decreased with time in an SBS concentration-dependent manner. However, this incompatibility may be clinically acceptable when the final SBS concentration is lower than 0.1 mg/mL and the mixed solution is used within 7 days. The results of the present study also indicate that the SBS concentration, as well as its presence/absence, should be considered when assessing the appropriateness of mixing OCT with other drugs. Furthermore, it may be necessary to examine the influences of the initial concentration of OCT and concentrated dextrose in order to ensure its stability when it is mixed with other drugs.

What this paper adds

What is already known on this subject

▸ When aggressive treatment is not an option, octreotide (OCT) is an important drug that can be used to maintain the quality of life of patients until the end stage.

▸ Large variations have been reported in the efficacy of OCT (44%–100%).

▸ OCT is a cyclic octapeptide whose hydrolysis is promoted by sodium bisulfate (SBS). However, the critical SBS concentration is unknown.

What this study adds

▸ We examined reproducibility, with a focus on the SBS concentration-dependent OCT stability of drugs made from morphine and metoclopramide, using high performance liquid chromatography.

▸ This study clarified the critical concentration of SBS for OCT stability.

Competing interests None.

Patient consent Obtained.

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