Effect of early oral nutrition supplement using Encover in patients undergoing hepato-biliary-pancreatic surgery

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Backgrounds/Aims: Early recovery after surgery has become a popular trend. The aim of this study was to evaluate effect of nutritional intervention using Encover, an oral nutritional supplement, in patients undergoing hepato-biliary-pancreatic surgery.

Methods: This single center, prospective case-control study was conducted in Gangnam Severance Hospital from September 2018 to April 2019. Through randomization, patients were divided into an experimental group (30 patients) and a control group (30 patients). At postoperative seven days, the experimental group was instructed to take two packs of Encover (JW Pharmaceutical, Seoul, Korea) daily for seven days. Body cell mass index was measured at seven days after surgery and 14 days after discharge and Patient-Generated Subjective Global Assessment (PG-SGA) was performed at 14 days after discharge.

Results: Body cell mass index during outpatient follow-up was significantly decreased compared to that at discharge in both groups. However, the amount of body cell mass index showed no significant difference between postoperative seven days and outpatient follow-up in either group. During outpatient follow-up, the experimental group had a higher mean value of PG-SGA score than the control group (11.32 ± 3.46 vs. 9.48 ± 3.97; p = 0.037).

Conclusions: Short-term Encover doses after surgery may not produce significant results in weight gain or other body cell mass index. Encover did not significantly affect other dietary conditions based on PG-SGA.

Key Words: Nutritional support; Supplementary feeding; Dietary supplements; Enhanced recovery after surgery; Enteral nutrition

INTRODUCTION

It has been reported that 20% to 50% of hospitalized patients are malnourished, with nutritional status worsening during hospitalization [1-4]. Poor dietary intake during hospitalization can cause deterioration of nutritional status. Malnutrition can increase complications including infection. It can also increase hospital stay and mortality [3,5,6]. Thus, appropriate nutritional therapy is needed. It is essential to point out areas to raise awareness for medical staff. In particular, in the case of gastrointestinal cancer, catabolism increases up to 10 days after surgery while protein anabolic activity decreases, which can result in loss of intestinal and skeletal muscle proteins in the body, postoperative weight loss, and cachexia, leading to decreased ability to recover for the body [4,7].

Recent surgical trends are focusing on early recovery after surgery. Enhanced Recovery After Surgery has been introduced. Early nutrition is strongly recommended for rapid recovery and proper nutrition of surgical patients [8-10]. However, when having a liquid to soft food diet after surgery, it is difficult to meet nutritional requirements due to the low caloric content per unit volume. After surgery, nutritional adequacy of a patient should be carefully reviewed due to problems such as indigestion, early postprandial fullness, bloating, and restriction of one-time meal intake [11,12]. Therefore, it is important to provide patients with a diet having proper amounts and adequate calories for sufficient nutrition to help their recovery.

Encover (JW Pharmaceutical, Seoul, Korea) is a product developed for patients who have difficulty in eating or lack nutritional intake [13,14]. It is an enteral nutrient that is used for tuberous nutrition, especially when oral nutrition is difficult for a long period of time. Patients who undergo major hepato-bil-
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Biliary-pancreatic surgery have a relatively low survival rate. Low body weight (body mass index [BMI] < 18.5 kg/m²) is directly related to the factors that increase the 5-year survival rate, which is less than 20% [15]. Intake of Encover is expected to improve body weight and biochemical/human measurements by contributing to continuous nutritional supplementation and increasing energy and protein intake. It is thought that it can increase survival rate by improving the quality of life of patients with liver and gallbladder disease.

The purpose of this study was to evaluate the effect of nutritional supplementation using Encover, an oral nutritional supplement, in patients undergoing major hepato-biliary-pancreatic surgery. Changes in weight, body fat, and muscle mass were determined after additional Encover was taken after surgery.

MATERIALS AND METHODS

Study design and period
This was a single center, randomized case control study. It was conducted from September 2018 to April 2019. Patients in the Hepatobiliopancreatic Cancer Clinic, Gangnam Severance Hospital, Yonsei University College of Medicine were recruited. This study was approved by Gangnam Severance Institutional Review Board (approval number: 3-2017-0222). Written informed consent was obtained from all participants.

Patient selection and enrollment criteria

Inclusion criteria
- Patients scheduled for major hepato-biliary-pancreatic surgery
- Major hepato-biliary-pancreatic surgery: segmentectomy, hemihepatectomy, segmental resection of bile duct, radical cholecystectomy, distal pancreatectomy, pancreatoduodenectomy

Exclusion criteria
- Patient with poor adherence to oral nutritional supplements
- Liver failure or renal failure patients
- Patient unable to intake food orally
- Patient with severe ascites and edema affecting weight evaluation
- Patient whose cancer has metastasized to the brain

Fig. 1. Patient-Generated Subjective Global Assessment sheet.

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• Patient who have difficulty controlling blood sugar
• Patient with BMI > 30 kg/m²
• Illiteracy/foreign patient
• Patient with contraindications for Encover administration

Sample size calculation
The output of the sample size was based on an independent two-sample t-test. The expected ratio of maintaining muscle mass was determined by referring to a paper previously published by Kim et al. [16] Assuming an alpha value of 0.05 and 1-β (power) of 0.8, 22.271 samples would be needed for each group. Considering a dropout rate of 30%, 30 samples would be needed for each group.

Perioperative evaluation and intervention
All patients who were potential candidates for major hepatobiliary-pancreatic surgery were informed about this study. Only patients who voluntarily consented participated in this study. Patients who agreed to participate in this study were assigned to either a control group or an experimental group by randomization. Randomization was take place via an allocation randomization system before surgery. Patients were randomized to one of the two groups at a 1 : 1 ratio.

After patients were enrolled, their baseline characteristics were recorded. Patient-Generated Subjective Global Assessment (PG-SGA) (Fig. 1) was used to evaluate and record their current nutritional status. Body cell mass index measurements (Inbody S-10; Biospace, Seoul, Korea) were taken before surgery. At postoperative seven days, body cell mass index measurement was taken again. Patients were excluded from this study if they could not tolerate soft diet and/or refused to take Encover on postoperative seven days. At 14 days after discharge, the last body cell mass index measurement was taken. PG-SGA was performed during outpatient follow-up. Patients taking less than 7 packs in total were excluded from this study. During the study period, a total of 8 cases were excluded from the Encover group as study denied or under dose in the experimental group. A total of 3 cases withdrew from this study due to operation hold, study denied, or complications in the control group (Fig. 2).

Statistical analysis
All statistical analyses were performed using IBM SPSS software, ver. 25.0 (IBM Corp., Armonk, NY, USA). Categorical variables were analyzed either by chi-squared test or Fisher’s exact test, while continuous variables were analyzed using Student’s t-test or Mann–Whitney U test. Statistical significance was considered when p-value was less than 0.05.

RESULTS
Clinicopathologic characteristics of patients are presented in Table 1. There was no significant difference in sex, age, or perioperative laboratory data related to nutritional status (such as albumin, prealbumin, cholesterol) between the control group and the experimental group. Major diagnosis, open and laparoscopic ratio, and main operation site (liver or pancreas) did not differ between the two groups either. Preoperative body cell mass index did not show significant difference between the two groups either (Table 2). During outpatient follow-up, body weight, body cell mass, soft lean mass, and fat free mass, but not fat mass, were significantly decreased than those at postoperative seven days in both groups (Table 3). When comparing the amount of change in body cell mass index from postoperative seven days to outpatient follow-up, there was no difference between the two groups. Body weight was decreased by 3.82 ± 2.84 kg in the control group and 4.27 ± 3.65 kg in the experimental group, showing no significant difference between the two (p = 0.627). There was no significant difference in the amount of change in body cell mass (p = 0.684), soft lean mass

![Fig. 2. Case enrollment diagram.](https://doi.org/10.14701/ahbps.21-152)
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When PG-SGA score and grade were compared, there was no difference in preoperative PG-SGA score or grade between the two groups. However, during postoperative outpatient follow-up, the experimental group had higher mean PG-SGA score (p = 0.037) and PG-SGA grade (p = 0.032) than the control group (Table 5).

DISCUSSION

This study was conducted in patients who underwent hepatobiliary-pancreatic surgery. The purpose of this study was to determine the effect of nutritional supplement using Encover by comparing postoperative body cell mass index following administration of additional oral nutritional supplements.

In this study, body cell mass index analysis was performed using the multi-frequency impedance method with Inbody S-10. This method has been used in many studies as a simple and effective method to indirectly measure the body cell mass index [17-19]. PG-SGA is also widely used as a tool to evaluate the nutritional status of patients by examining changes in body weight, changes in meal intake, problems related to meals, and the level of physical activity through interviews with experi-

### Table 2. Preoperative body cell mass index difference between the experimental group and the control group

| Variable            | Control group (27 cases) | Experimental group (22 cases) | p-value |
|---------------------|--------------------------|-----------------------------|---------|
| Body weight (kg)    | 64.75 ± 12.13            | 66.03 ± 12.01               | 0.715   |
| Body cell mass (kg) | 32.02 ± 7.80             | 32.77 ± 5.37                | 0.693   |
| Soft lean mass (kg) | 46.30 ± 10.78            | 47.50 ± 7.66                | 0.660   |
| Fat free mass (kg)  | 49.20 ± 11.37            | 50.40 ± 8.06                | 0.677   |
| Fat mass (kg)       | 15.56 ± 7.69             | 15.62 ± 7.20                | 0.975   |

Values are presented as mean ± standard deviation.

### Table 3. Body cell mass index at postoperative seven days and outpatient follow-up

| Variable            | Discharge | OPD F/U | p-value |
|---------------------|-----------|---------|---------|
| Body weight (kg)    | 65.8 ± 11.50 | 62.0 ± 11.36 | < 0.001 |
| Control group (27 cases) | 66.6 ± 11.18 | 62.3 ± 9.67 | < 0.001 |
| Experimental group (22 cases) | 32.2 ± 7.07 | 30.1 ± 6.67 | < 0.001 |
| Body cell mass (kg) | 32.7 ± 5.21 | 30.2 ± 5.04 | < 0.001 |
| Control group (27 cases) | 47.6 ± 9.72 | 44.1 ± 9.15 | < 0.001 |
| Experimental group (22 cases) | 48.3 ± 7.69 | 44.2 ± 7.30 | < 0.001 |
| Fat free mass (kg)  | 50.7 ± 10.22 | 47.0 ± 9.59 | < 0.001 |
| Control group (27 cases) | 51.4 ± 8.11 | 47.0 ± 7.68 | < 0.001 |
| Experimental group (22 cases) | 15.1 ± 7.60 | 15.1 ± 7.14 | 0.900   |
| Fat mass (kg)       | 15.2 ± 7.29 | 15.3 ± 6.40 | 0.870   |

Values are presented as mean ± standard deviation.

OPD F/U, outpatient follow-up.
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Table 4. Change amount in body cell mass index between postoperative seven days and outpatient follow-up

| Variable                  | Change amount | p-value |
|---------------------------|---------------|---------|
| Body weight (kg)          | −3.82 ± 2.84  | 0.627   |
| Control group (27 cases)  | −4.27 ± 3.65  |         |
| Experimental group (22 cases) | −2.19 ± 2.43  | 0.684   |
| Control group (27 cases)  | −2.45 ± 1.92  |         |
| Experimental group (22 cases) | −4.06 ± 3.16  | 0.561   |
| Soft lean mass (kg)       | −3.46 ± 3.83  |         |
| Control group (27 cases)  | −4.36 ± 3.36  |         |
| Experimental group (22 cases) | −2.45 ± 1.92  | 0.578   |
| Fat free mass (kg)        | 3.75 ± 4.07   | 0.834   |
| Control group (27 cases)  | −4.36 ± 3.36  |         |
| Experimental group (22 cases) | −0.07 ± 2.64  |         |
| Fat mass (kg)             | 0.09 ± 2.39   |         |
| Control group (27 cases)  | −2.19 ± 2.43  |         |
| Experimental group (22 cases) | −2.45 ± 1.92  |         |

Values are presented as mean ± standard deviation.

In conclusion, short-term Encover doses after surgery may not produce significant results in weight gain or other body cell mass index. In addition, PG-SGA score and grade at outpatient follow-up were higher than those in the control group, indicating malnutrition in the experimental group.

Previous study have hypothesized that insufficient oral food intake is correlated with satiety and volume in the digestive tract [16]. Therefore, oral nutritional supplements such as high energy density in the diet are expected to help increase final caloric intake and subjective nutritional indicators after ingestion. However, PG-SGA score and grade showed opposite results. Taking Encover two packs a day might have reduced their original normal meal intake due to worsening of gastrointestinal satiety. This is a short-term side effect, suggesting that additional oral nutritional supplements may affect dietary intake. Additional oral nutritional supplements cannot be free from risks such as reduced intake of regular meals that can be better absorbed. In the future, in terms of nutritional management, when taking alternative nutrients, it is necessary to continuously monitor patient’s subjective nutritional status and closely monitor whether there is a possibility of digestive disorders or reduced intake.

This study has some limitations. First, it did not have a long-term follow-up after surgery. Interference aspects such as uncontrolled variables (gastrointestinal trouble) should also be considered. The effect on energy consumption rate according to the difference in exercise amount after each patient’s operation cannot be completely excluded.

In future studies, rather than using a short one-week dose period, we intend to investigate difference in body cell mass index according to the increase or decrease in dose by considering expansion and consumption of Encover as continuous variables.

In conclusion, short-term Encover doses after surgery may not produce significant results in weight gain or other body cell mass index. In addition, Encover does not significantly affect other dietary conditions based on PG-SGA.

https://doi.org/10.14701/ahbps.21-152
ACKNOWLEDGEMENTS
This study was an investigator-initiated trial supported by JW pharmaceutical.

FUNDING
None.

CONFLICT OF INTEREST
No potential conflict of interest relevant to this article was reported.

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