A single-day mouse mesenteric lymph surgery in mice: an updated approach to study dietary lipid absorption, chylomicron secretion, and lymphocyte dynamics

Nikolaos Dedousis, Lihong Teng, Jitendra S. Kanshana, and Alison B. Kohan

Department of Medicine, Division of Endocrinology and Metabolism, University of Pittsburgh, School of Medicine, Pittsburgh, PA, USA

Abstract The intestine plays a crucial role in regulating whole-body lipid metabolism through its unique function of absorbing dietary fat. In the small intestine, absorptive epithelial cells emulsify hydrophobic dietary triglycerides (TAGs) prior to secreting them into mesenteric lymphatic vessels as chylomicrons. Except for short- and medium-chain fatty acids, which are directly absorbed from the intestinal lumen into portal vasculature, the only way for an animal to absorb dietary TAG is through this chylomicron/mesenteric lymphatic pathway. Isolating intestinal lipoproteins, including chylomicrons, is extremely difficult in vivo because of the dilution of postprandial lymph in the peripheral blood. In addition, once postprandial lymph enters the circulation, chylomicron TAGs are rapidly hydrolyzed. To enhance isolation of large quantities of pure postprandial chylomicrons, we have modified the Tso group’s highly reproducible gold-standard double-cannulation technique in rats to enable single-day surgery and lymph collection in mice. Our technique has a significantly higher survival rate than the traditional 2-day surgical model and allows for the collection of greater than 400 μl of chylous lymph with high postprandial TAG concentrations. Using this approach, we show that after an intraduodenal lipid bolus, the mesenteric lymph contains naïve CD4+ T-cell populations that can be quantified by flow cytometry. In conclusion, this experimental approach represents a quantitative tool for determining dietary lipid absorption, intestinal lipoprotein dynamics, and mesenteric immunity. Our model may also be a powerful tool for studies of antigens, the microbiome, pharmacokinetics, and dietary compound absorption.

Supplementary key words small intestine • mesenteric lymph • lipoproteins • metabolism • dietary fat • enterocytes • fatty acid/transport • lymphocytes • triglycerides • CD4+ T cells • postprandial lipids

The intestine plays a crucial physiological role in whole-body lipid metabolism through dietary fat absorption. In the small intestine, absorptive epithelial cells emulsify hydrophobic dietary triglycerides (TAGs) prior to secreting them into the mesenteric lymphatics as chylomicron particles (1). Dietary TAG first enters the duodenum and is hydrolyzed by pancreatic lipase in the small intestinal lumen. Free fatty acids and monoglycerol, which are hydrolysis products of dietary TAG, are then absorbed into the enterocyte, resynthesized into TAG, and packaged into chylomicrons (reviewed here (2, 3)). Mature chylomicrons are then secreted through the basolateral membrane into the mesenteric lymphatics (4). Except for short- and medium-chain TAG, which are directly absorbed from the intestinal lumen into portal blood, the only way for an animal to absorb dietary TAG is through this chylomicron/mesenteric lymphatic pathway (5, 6).

The human small intestine is almost always secreting lipoproteins. These include intestinal HDLs, a major pathway for cholesterol absorption; intestinal VLDL that can contribute up to ~11% of the TAG in plasma; and the major intestinal lipoprotein, the chylomicron, which is the major carrier of dietary TAG (7–9). Chylomicrons are secreted as early as 13 min after a lipid meal and reach their peak secretion rate at ~2 h after a meal (10). Chylomicron secretion continues over the next ~6 h, which is considered the postprandial period (1). Several studies have demonstrated that oral glucose can also mobilize stored intracellular TAG in enterocytes and can potentiate the secretion of lipid-poor chylomicrons for up to 16 h after the last meal (12, 13).

While chylomicron assembly is largely driven by TAG in the diet, there are situations where enterocyte de novo lipogenesis can also provide fatty acids for intestinal lipoproteins (14). An important example is during the fasting state, when apical fatty acids are absent, yet the small intestine still secretes apoB48-containing particles (essentially a constitutive intestinal VLDL) (8, 15, 16). In addition, there is evidence that diet-induced insulin resistance and hypertriglyceridemia results in the overproduction of both chylomicrons and intestinal VLDL, and that these
particles are synthesized and plasma fatty acids (17–19). Therefore, by the time most people are ready to eat their next meal, they are still in the postprandial state of the previous meal, and even when they enter a fasting period, the small intestine still secretes apoB-containing particles with TAG and cholesterol.

Diseases where dietary fat absorption is defective include inflammatory bowel disease, cystic fibrosis, short bowel syndrome, intestinal wasting diseases (20, 21), and lysosomal acid lipase deficiencies (like Wolman disease or cholesterol ester storage disease) (22–25). Defective dietary fat absorption is not simply an inability to extract fat from a meal (since the gut can adapt to defects in surface area and length), it also results in the use of noncanonical absorption pathways that form abnormal and potentially dysfunctional chylomicrons. There are many molecular events that have dramatic effects on chylomicron secretion; these include intracellular dysfunction in dietary TAG absorption and intracellular resynthesis (26–29), endoplasmic reticulum events like the lipidation of nascent chylomicrons (30, 31), and changes in the source of chylomicron TAG (i.e., from dietary fatty acids, or less frequently, from de novo fatty acids and glycerol phosphate) (32, 33). The net result is often a rescue of total fat absorption but at the cost of normal chylomicron synthesis and secretion.

A key component of chylomicrons are their apolipoproteins, which regulate multiple metabolic events. ApoB-48 is the structural protein making up chylomicrons (30, 34–36). ApoC-III and apoC-II work in opposition to each other to tightly regulate the clearance of chylomicron TAG from blood into tissues (apoC-III inhibits LPL hydrolysis (37) and LDL receptor-mediated endocytosis (38), respectively, and apoC-II stimulates LPL activity (39)). Chylomicrons acquire apoE in the plasma, which is required for the ultimate clearance of remnant chylomicrons from blood into the liver (40). Together, these activities of apolipoproteins are crucial to ensuring not only that extraintestinal tissues have access to dietary TAG but also that TAG is also efficiently removed from the circulation into the liver.

When apolipoproteins are missing or defective, dietary TAG is not metabolized normally. For example, chylomicrons containing excess apoC-III inhibit LPL and LDL receptor clearance pathways so successfully that these particles are retained in plasma and cause severe hyperlipidemia (41–44). Interestingly, excess apoC-II is also associated with reduced LPL activity and hypertriglyceridemia (45, 46). Clinically, dysfunction in the clearance of chylomicron TAG from blood is a major clinical problem (47–49). Children with short bowel disease, or patients missing small intestine, are often treated with parental intravenous lipids, which contain emulsified TAG but not apolipoproteins (48, 50). Though this effectively delivers soluble TAG to the blood (like chylomicrons do), the bulk of intravenous lipids is rapidly cleared by the liver, resulting in fatty liver and inefficient calorie replenishment (50, 51). This clinical scenario highlights the physiological importance of chylomicrons and the critical role of the intestine in the postprandial state.

Causes of abnormal chylomicron synthesis include: 1) inability to maintain the primary site of lipid absorption: The proximal duodenum and jejunum have the highest fat absorption and chylomicron synthesis capacity (52). If dietary fat is unabsorbed in these proximal locations, and instead travels to the ileum, the chylomicrons produced will often contain less TAG but more bacterial lipopolysaccharide (LPS) per particle (53, 54). Bacterial LPS is known to travel through enterocytes, and during this process, it can be incorporated on chylomicrons. The amount of LPS can be temporarily reduced when chylomicron secretion is blocked with Pluronic L81 (55), but the dynamics of this process are not clear. 2) Defects in the trafficking of dietary TAG through absorptive enterocytes: there are three regulated steps of dietary lipid absorption—luminal hydrolysis and fatty acid traffic through the unstirred water layer into the mucosa, intracellular fatty acid re-esterification into TAG, and enterocyte chylomicron secretion into lymph. Dysfunction at each step yields small dense chylomicrons or chylomicrons containing inappropriate apoproteins. These changes can lead to either inefficient metabolism of chylomicrons by lipases on peripheral tissues and chylomicron retention in the blood (56, 57), or alternatively, can cause chylomicrons to speed through the vasculature avoiding tissue lipid delivery and instead load the liver with dietary fat (as in chylomicrons lacking apoC-III or containing excess apoC-II) (58–60). This deprives peripheral tissues of metabolic substrates (61). These processes are an underappreciated mechanism in how the small intestine can direct whole-body metabolism.

Because of these complexities in chylomicron character, metabolism, and formation, it is difficult to find a direct replacement or comparator for mechanistic studies (62, 63). Our group and many others routinely use fatty acids bound to albumin or lipid emulsions because they are so much simpler to prepare and dose than chylomicrons (64–68). Isolating intestinal lipoproteins is not trivial. Chylomicron remnants can be isolated from plasma via ultracentrifugation in the 2–6 h after a gavage, but these particles will be a mixture of partially hydrolyzed chylomicrons varying in size and TAG content (69). It is also difficult to isolate a large quantity of chylomicron remnants from plasma, especially in quantities needed for cell culture experiments. Another approach is to cannulate the thoracic lymph duct prior to its intersection with the subclavian vein, which will capture mesenteric lymph as it enters the circulation (70). Again, large quantities of lymph are difficult to obtain using this approach. Finally, lymph can be statically sampled from the mesenteric lymph duct at a single time point after a lipid meal (71, 72).
This approach requires perfusion of the gut but does not allow for terminal measurement of the movement of dietary lipids through the gastrointestinal tract or their absorption capacity between the lumen and enterocytes. Lymph volumes are also quite small.

It should also be emphasized that intestinal lymphatic system is not merely a passive duct for drainage of fats and for fluid balance. Lymph is an immune compartment. In addition to absorbing dietary TAG and other nutrients, the intestine must also regulate immune homeostasis in response to intestinal microbiota while maintaining vigilance against infectious microorganisms. This task is complex; microbiota contain a multitude of antigens, expressed on organisms that have heterogeneous interactions with the host and are constantly changing both in composition and behavior. In addition, the immune response must also remain sensitive to inflammatory signals produced by damaged or infected enterocytes. There is a major interest in the dynamics of inflammatory versus suppressive lymphocytes in the small intestine, and the mesenteric lymph (both postnodal and prenodal compartments) is a major site of these dynamics (72–74).

Using the Tso Lab’s highly reproducible gold-standard double-cannulation technique as the foundation for our studies, we now describe a 1-day surgical model for isolating mesenteric lymph. Our procedure differs from previous design by reducing the surgery and experimental period to a single day, improving TAG excursion in response to duodenal lipids, and dramatically improving surgical success rate, which reduces the number of experimental animals needed.

**MATERIALS AND METHODS**

**Animals**

C57BL/6J mice were obtained from Jackson Labs (Bar Harbor, ME), and in-house bred WT mice aged 8–14 weeks were used. All mice were housed on a 12-h light/dark cycle with ad libitum access to standard chow and water. Whether mice are fasted overnight or not will depend on the experimental design. In practice, we find that an overnight fast is not necessary unless we are concerned about differences in rate of stomach emptying. For all experiments presented here, mice were not fasted overnight prior to surgery. All surgical procedures were approved by the University of Pittsburgh Internal Animal Care and Use Committee (protocol #20047008) and comply with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

**Cannulation of mesenteric lymph and intraduodenal infusion tube**

For these procedures, mice are placed under isoflurane anesthesia (induction at 5% and 2% maintenance during surgery), and both a mesenteric lymph duct cannula and duodenal infusion tube are installed. Prior to surgery, the mice are placed on a heated surgical pad. A surgical incision is made along the midline, and the abdominal viscera were manipulated with a retractor to expose the mesenteric lymph duct. The duct is partially cut at the proximal end, and the catheter tip (micro-Renathane tubing cut on the bias to form a cannula tip; Braintree Scientific; catalog no: MRE-095) was inserted at the outermost part of the duct and fastened with a drop of Krazy glue. For placement of the duodenal feeding tube, the stomach is exposed and drawn partially out of the abdominal cavity. We use an 18G needle to insert the micro-Renathane tubing (Braintree Scientific; catalog no: MRE-037) into the stomach and just past the pyloric sphincter into the duodenum. This is secured with a purse-string suture. Using 5-0 suture, we closed the incision and placed the mouse in Snuggle restraint jacket (mouse Snuggle; Lomir, Inc, https://maps.google.com/?q=213+West+Main+Street+Malone,+NY+12953&entry=gmail&source=; Malone, NY; MS02.5PM). Snuggled mice are placed on a rotator table in a temperature and humidity-controlled polycrystalline filter-top container (12” W × 14” L × 8” H). The externalized duodenal feeding tube is connected to a Harvard infusion pump. The externalized lymph catheter is carefully placed to allow gravity flow of lymph into Eppendorf collection tubes on ice. Mice stay in the chamber throughout the experiment. Postsurgery, mice continuously receive 5% glucose in saline, at a 0.3 ml/h infusion rate, via duodenal tube to compensate for fluid and electrolyte loss because of lymphatic drainage. The original Tso Lab 2-day lymph fistula model used postsurgery and overnight infusion of glucose/saline infusion to keep rats well hydrated (they cannot resorb the fluid lost in diverted lymph) (75). Because the mesenteric lymph is diverted, animals cannot be calorically replenished with lipids overnight; so instead glucose is preferable. Fluids must be infused via the intraduodenal infusion tube because the animals are restrained and cannot access water. In the Tso Lab protocol, rats are restrained in Bollman cages; in our protocol, mice are restrained with Snuggles. Either way, if the restraint is loosened enough for ad lib access to water, the animals will chew out their stitches and cannulas.

**Lipid infusion and collection of hourly lymph**

Mice are given a 300 μl bolus (which takes 2–3 min to infuse the total volume), of lipid (SMOFlipid 20% lipid injectable emulsion; Fresenius Kabi AG) followed by continuous glucose/saline at 0.3 ml/h.

**Choice of infusion TAG and amount.** Each 100 ml of SMOFlipid 20% contains refined soybean oil (60 g), medium-chain TAG (60 g), refined olive oil (50 g), fish oil (30 g), purified egg phospholipids (12 g), all-rac-α-tocopherol (16–23 mg), glycerol (25 g), sodium oleate (30 mg), and sodium hydroxide to adjust pH. This test lipid is designed for clinical use in patients receiving intravenous parenteral nutrition. It contains lipids that are traditionally absorbed via both the lymphatic and portal route (refined soybean and olive oils and medium-chain TAG, respectively) (76). Lymph was collected on ice for 1 h prior to lipid infusion (which represents a “fasting” experimental time point) and then hourly for 6 h continuously after the bolus infusion. Weight of each hourly lymph samples was recorded and used for flow rate.

**1-Day protocol.** One hour prior (~1) to the bolus lipid infusion, mice received a continuous duodenal infusion of 5% glucose in sterile saline, and a single hour of “fasting” lymph is collected. At time 0, the continuous intraduodenal infusion is switched to a bolus infusion of 0.3 ml SMOFlipid (which
takes ~2–3 min to infuse). Immediately after the bolus lipid, the continuous glucose infusion is resumed. Lymph is then collected hourly on ice for 6 h after the bolus infusion. At 6 h, mice are euthanized, and tissues are collected.

2-Day protocol. In parallel with the 1-day mice, 2-day mice receive a continuous duodenal infusion of 5% glucose in sterile saline after the surgical implantation of both cannulas. The design branches at time 0, which would be the time for bolus lipid infusion in a 1-day model. Instead, 2-day mice continue to receive the continuous duodenal infusion of 5% glucose in sterile saline for the next ~18 h (through the overnight period). On the morning of the second day, a single hour of “fasting” lymph is collected. At this point, as in the 1-day model, at time 0, the continuous intraduodenal infusion is switched to a bolus infusion of 0.3 ml SMOFlipid. Immediately after the bolus lipid, the continuous glucose infusion is resumed. Lymph is then collected hourly on ice for 6 h after the bolus infusion. At 6 h, mice are euthanized, and tissues are collected.

Flow cytometry

Cells from lymph collected prelipid and postlipid infusion were stained with antibodies on ice and protected from light. Intracellular staining was performed using the eBioscience Foxp3 staining kit as per the manufacturer's instructions (Thermo Fisher Scientific; catalog no.: TR210). Briefly, 2 μl of the diluted samples was incubated with 200 μl of enzyme reagent at 37°C for 5 min in a 96-well plate. The plate was read by MULTISKAN G0 (Thermo Fisher Scientific, Waltham, MA) plate reader at 500 nm. The absence of the dye indicates live cells. Live cells were then identified by leukocytes using CD45 (BioLegend, San Diego, CA; catalog no.: 103106) by FSC-A and then further identified as helper T cells with CD4 (Thermo Fisher Scientific; catalog no.: 45-0042-82) by FSC-A. To distinguish naïve CD4 cells, CD4 was gated by CD25+CD44lowCD62hi (Thermo Fisher Scientific; catalog nos.: 25-0251-82, 48-0441-82, and 17-0621-82, respectively), and for Treg cells, CD4+ was gated by CD25+ Foxp3+ as double-positive cells. Fluorescence minus one controls were used to determine all gates except FSC by side scatter and single cells. When appropriate, Fc receptors were blocked using CD16/CD32 Monoclonal Antibody (Thermo Fisher Scientific; catalog no: 14-0161-81), before staining with antimouse fluorochrome-conjugated antibodies FlowJo software (Treestar, Ashland, OR) was utilized for final analysis of collected data. Thermo Fisher Scientific and BioLegend antibodies used are described in Table 1.

TABLE 1. Reagents used for flow cytometry analysis

| Reagent | Catalog number | Source |
|---------|----------------|--------|
| CD4 monoclonal antibody (RM4-5), PerCP-Cyanine5.5, eBioscience™ (1:100 dilution) | 45-0042-82 | Thermo Fisher Scientific |
| CD25 monoclonal antibody (PG6.15), PE-Cyanine7 (1:250 dilution) | 25-0251-82 | Thermo Fisher Scientific |
| CD44 monoclonal antibody (IM7), eFluor 450, eBioscience™ (1:100 dilution) | 48-0441-82 | Thermo Fisher Scientific |
| CD62L (L-Selectin) Monoclonal Antibody (ME1-11), APC (1:400 dilution) | 17-0621-82 | Thermo Fisher Scientific |
| FOX3 Monoclonal Antibody (FJK-16s), Alexa Fluor 488 (1:200 dilution) | 55-5773-82 | Thermo Fisher Scientific |
| LIVE/DEAD™ Fixable Near-IR Dead Cell Stain Kit | L10119 | Thermo Fisher Scientific |
| eBioscience™ Foxp3/Transcription Factor Staining Buffer Set | 00-5523-00 | Thermo Fisher Scientific |
| eBioscience™ Flow Cytometry Staining Buffer | 00-4222-26 | Thermo Fisher Scientific |
| PE anti-mouse CD45 Antibody (1:250 dilution) | 14-0161-81 | Thermo Fisher Scientific |
| CD16/CD32 Monoclonal Antibody, eBioscience™ (1:100 dilution) | 00-4222-26 | Thermo Fisher Scientific |

Statistics

Values are expressed as mean ± SEM. The differences between two groups were analyzed by Student's t test, and the lymph flow rate and lymphatic TAG concentrations were analyzed by multiple unpaired t tests using GraphPad Prism 9 (GraphPad Software). Differences were considered statistically significant at P < 0.05.

RESULTS

Adapted the classic murine conscious lymph fistula model (6, 77–79) to a single day experimental design

In our 1-day surgical model, the implantation of double cannulas is immediately followed by duodenal lipid infusion for the delivery of dietary TAG directly to the small intestinal lumen (bypassing stomach emptying, pancreatic enzyme secretion, or bicarbonate buffering dysfunction that can occur in diseases of the small intestine (21, 80)).

The 1-day surgical model is summarized in Fig. 1. Double cannulation takes approximately 2–4 h. Mice recover in Snuggle restraints and in a warmed and humidified incubator. One hour prior to the bolus lipid infusion, mice receive a continuous duodenal infusion of 5% glucose in sterile saline, and a single hour of “fasting” lymph is collected. The continuous scatter area, followed by identification of single cells using FSC height by FSC width. Finally, live cells were gated on using FSC-A by Live/Dead viability dye (Near-IR Dead Cell Stain Kit; Thermo Fisher Scientific; catalog no: L10119).
intraduodenal infusion is then switched to a bolus infusion of 0.3 ml SMOFlipid. Immediately after the bolus lipid, the continuous glucose infusion is resumed. Lymph is collected on ice for 6 h continuously at 1-h intervals after the bolus infusion (for a total of six samples, each 1 h of lymph flow).

Figure 2 shows anatomical models and surgical photos of the 1-day technique.

This differs from the classic 2-day lymph fistula model by several points: 1) 2-day surgical model mice receive a continuous duodenal infusion of 5% glucose in sterile saline and a single hour of “fasting” lymph is collected. At time 0, the continuous intraduodenal infusion is switched to a bolus infusion of 0.3 ml SMOFlipid. Immediately after the bolus lipid, the continuous glucose infusion is resumed. Lymph is collected hourly on ice for 6 h after the bolus infusion. At 6 h, mice are euthanized, and tissues were collected. C: Two-day experimental model. Mice receive a continuous duodenal infusion of 5% glucose in sterile saline as in (B), but the infusion continues for an additional ~18 h. D: On the morning of the second day, a single hour of “fasting” lymph is collected. As in (B), at time 0, the continuous intraduodenal infusion is switched to a bolus infusion of 0.3 ml SMOFlipid. Immediately after the bolus lipid, the continuous glucose infusion is resumed. Lymph is collected hourly on ice for 6 h after the bolus infusion. At 6 h, mice are euthanized, and tissues were collected. Created with BioRender.com.

**Improvement survival rates after 1-day surgical model design**

Surgical success is considered a mouse that has milky lipid-rich flowing lymph starting ~15 min after the duodenal lipid infusion and lasting at least 5 h. If an animal died prior to the start of the lipid infusion, or had no lymph flow after duodenal infusion, this qualifies as an experimental loss. We found that animals that underwent the 1-day surgical design had a 63.0% success rate, with 17 of 27 mice (Fig. 3A). In the classic 2-day surgical design, mice received the glucose/saline infusion immediately postsurgery and throughout an overnight recovery period. These mice received the duodenal lipid infusion on the morning of the second day. These mice were much more likely to die during the recovery period prior to the duodenal infusion, and they were more likely to lose steady lymph flow during either the glucose/saline infusion or in response to duodenal lipid. The success rate for this surgical design was 38.0% (8 of 21 mice), which is much lower than the 1-day model.

**Comparison in lymph flow rates and total lymph secretion between 1-day and 2-day experimental models**

We collected a 1 h of lymph flow from each mouse in the hour prior to duodenal lipid infusion (mice...
received the infusion of glucose/saline except during the 2–3 min lipid infusion, and then we collected hourly lymph samples for 6 h and kept lymph on ice. We found that the basal flow rate was not different between the two groups (Fig. 3B, time 0). Flow rates for both groups increased from 1 to 3 h after lipid infusion at the same rate. In rats, this increased lymph flow is a vagal response to duodenal lipid (81). Our data are the first to suggest that this occurs in mice as well as rats. Lymph flow rates only differ in the late postprandial period at hour 5 between groups (Fig. 3B, time 5). Overall, there is not a significant difference in the total amount (or volume) of lymph collected in 6 h between groups (Fig. 3C). This suggests that if mice survive the overnight recovery period, they can produce lymph volumes like their 1-day surgical model counterparts.

Increased TAG secretion in lymph in response to shortened experimental time

We measured lymphatic TAG concentrations following a bolus intraduodenal lipid infusion (Fig. 4A). In both groups, TAG secretion into lymph peaked at 3 h postlipid infusion. However, the magnitude of increase in TAG concentration is much higher in the 1-day compared with 2-day surgical model mice, and the 1-day mice secrete significantly more TAG into lymph than their 2-day counterparts over the entire course of lymph collection (Fig. 4B).

We wondered if the difference in TAG secretion could be attributed to the length of the recovery period because of the possibility that the long recovery period postsurgery diminishes small intestinal function. To test this possibility, we infused lipid and collected hourly lymph as in the 1-day design and followed this with an overnight recovery with continuous glucose/saline and a second lipid infusion on day 2 (experimental model depicted in Fig. 4D). Basal lymphatic TAG concentrations (0 h) were not different between the mice receiving their first or second lipid bolus, and after both bolus infusions, the lymphatic TAG peak at 3 h (Fig. 4C). Regardless of whether the mouse is receiving its second lipid bolus (as in Fig. 4C) or is receiving its first lipid bolus after an overnight recovery (as in first Fig. 4A), the magnitude of TAG in lymph is ~3× higher when the lipid infusion occurs on the first day than when it occurs on the second day. Therefore, the difference in TAG excursion is tied to length of time postsurgery, with 1-day lipid infusions resulting in higher lymphatic TAG concentrations.

Analysis of T lymphocytes in mesenteric lymph before and after a lipid bolus

Lymphocytes have been successfully isolated from ~5 μl static lymph and routinely from mesenteric lymph nodes (mLNs) but not from actively flowing lymph from mice receiving luminal nutrients (82–85).
Using our 1-day surgical model, mice received a continuous duodenal infusion of 5% glucose in sterile saline immediately after implantation of both cannulas, and then clear lymph was collected during this “pre-lipid bolus” period. At time 0, the continuous intraduodenal glucose/saline infusion is switched to a bolus infusion of 0.3 ml lipid. Immediately after the bolus of lipid, the continuous glucose infusion is resumed. “Postlipid bolus” lymph was then collected hourly on ice for 6 h continuously after the bolus infusion. Lymph was pooled from \( n = 2–3 \) mice during each experiment to have enough cells for flow cytometry analysis, and the experiment was repeated three times.

Lymph (\( \sim 0.5 \) ml) was resuspended in 5 ml PBS, pelleted, and all cells were then stained with a CD4\(^+\) T-cell panel. As shown in Fig. 5A, \( \sim 3–4 \) million cells were present in 1 ml of lymph. Of these, 98–100% were CD45\(^+\) leukocytes (Fig. 5B) and 40–46% CD4\(^+\) cells (Fig. 5C). Of the total population of CD4\(^+\) cells, 77–84% were naïve CD4 cells (Fig. 5D), which had \( \sim 3–4 \)% of CD4\(^+\)CD25\(^+\)Foxp3\(^+\) regulatory T cells (Fig. 5E). We found no significant differences between the populations of these cells in the prelipid versus postlipid bolus groups (summarized in Fig. 5E). This analysis demonstrates the presence of CD45\(^+\) leukocytes, CD4\(^+\) and naïve CD4 cells, and regulatory T cells, and the similar lymphocyte populations in the prelipid and postlipid bolus lymph. In addition, the 1-day lymph fistula surgical model is appropriate for quantitative fluorescence-activated cell sorting immune profiling in a variety of experimental conditions.

**DISCUSSION**

The mouse lymph fistula technique should be considered the gold-standard technique for determining the partitioning of dietary nutrients into lymph in response to duodenal nutrients. It has been used to successfully uncover major mechanisms including the role of apolipoproteins, intestinal lipoproteins, lipid absorption, the secretion of incretin hormones into lymph in response to both lipids and glucose, and the relative absorption of xenobiotic toxins through the portal and lymphatic routes (4, 78, 53, 86–90). Our contribution to this technique is to 1) reduce the experimental time from 2 days to 1 day; 2) improve the
magnitude of duodenally infused lipids in lymph; 3) increase the reproducibility of the approach in mice; and 4) improve animal welfare; and 5) reduce the number of mice needed because of the higher survival rate.

A major experimental hurdle for collecting mesenteric lymph is the number of mice needed to ensure that enough mice survive the surgery and recover prior to and after lipid infusion. This is especially critical when using precious genetically modified mice and their WT littermates. By increasing the survival rate, we reduce the costs associated with both cohort generation and surgical labor. Our results show that the 1-day surgical model not only reduces the number of mice needed in each cohort but also reduces the amount of time the mice must survive after surgery (from ∼24 to ∼6 h). It had been thought that the overnight recovery period would let the animal to recover postoperative gastric contraction, reduce any “hang-over” anesthesia effects on lymph flow, and move the experimental period away from the immediate postsurgery recovery period (Dr Patrick Tso, unpublished communications and (77)). Our experiments actually test this assumption, and we find that the 24-h recovery period is not necessary for lymph flow and may paradoxically make survival rates worse. This could be because recovery from major abdominal surgery cannot be achieved overnight and is fraught with risks of poor temperature regulation and hydration (in addition to the possibility that mice chew out their stitches or cannulas). Our updated surgical design reduces both unnecessary animal death and the potential for animal distress during this fragile postsurgical period. This supports a major goal of American Association for Accreditation of Laboratory Animal Care, which is animal “Replacement, Reduction, Refinement” (91).

We attribute increased survival not only to the reduction in surgical and experimental time but also to the use of alternative restraint after surgery. The lymph fistula technique has been primarily carried out in rats, which are restrained after surgery in Bollman restraint cages (70, 92). Both mice and rats move torsionally, even when their limbs are restrained, and they will rapidly and lethally chew out stitches and cannulas if not appropriated restrained. We have moved away from the Bollman restraint cage and instead used mouse Snuggles (which are essentially a soft restraint jacket) in both the 1-day and 2-day protocols described here. The Snuggles have two advantages: they keep the animals warm, and the mice are not

Fig. 4. Increased TAG secretion in lymph in response to shortened experimental time. A: In open circles, WT mice received intraduodenal lipid on the same day of surgery (1 day), \( n = 17 \). In black squares, WT mice received intraduodenal lipid after an overnight recovery period, on the day after surgery (2 days), \( n = 8 \). TAG concentrations in lymph in response to lipid bolus were determined by chemical assay. B: Total mass of TAG secreted during the 6 h after intraduodenal bolus. C: In white diamonds, WT mice received lipid on the same day as surgery (day 1). In black diamonds, the same mice received a second intraduodenal lipid dose the day after surgery— experimental model shown in (D). \( n = 10^+ \), and values are means ± SEM. \( * P < 0.05 \), \( ** P < 0.005 \), and \( *** P < 0.0001 \). Model created with BioRender.com.
able to move torsionally. We believe this approach improves quality of life in the hours postsurgery and improves survival after the surgery.

TAG appearance in lymph represents a direct measure of the capacity and flux of enterocyte chylomicron secretion (since there is no removal mechanism for chylomicrons in this compartment—no lipases and no tissue clearance mechanisms (33, 54)). The only way for chylomicrons to disappear from lymph is to enter the circulation at the thoracic duct. A fat tolerance test is often used as a proxy for dietary fat absorption capacity. If this is done in the presence of Poloxamer 407, it will block chylomicron metabolism and clearance by LPL and other lipid receptors in the circulation leading

Fig. 5. Analysis of T lymphocytes in mesenteric lymph before and after a lipid bolus. WT mice were fitted with mesenteric lymph cannula and intraduodenal feeding tube. Mice then received a continuous duodenal infusion of 5% glucose in sterile saline. During this “prelipid bolus” period, clear lymph was collected. At time 0, the continuous intraduodenal infusion is switched to a bolus infusion of 0.3 ml lipid. Immediately after the bolus lipid, the continuous glucose infusion is resumed. “Postlipid bolus” lymph was then collected hourly on ice for 6 h after the bolus infusion. A: Number of cells per volume (1 ml) of lymph fluid collected and quantification of percent of parent population. Data are pooled from three independent experiments (n = 2–3/experiment). B–E: Representative flow cytometric plots showing lymphocytes from equal volumes of prelipid or postlipid bolus lymph. Cells gated: Live lineage: negative for LIVE/DEAD stain, (B) total leukocyte; CD45+ cells gated on live cells, (C) CD4+ cell gated on CD45+ cells, (D) naive CD4 cells CD62hi CD44low, and (E) Treg cells CD25+ Foxp3+ gated on CD4+. Numbers on plots denote cell frequencies (in percentages) of indicated parent populations and SEM, respectively. Data are representative examples of three independent experiments with n = 2–3 mice per experiment.
to elevated plasma TAG coming from both liver and intestinal lipoproteins (93). This design cannot disentangle the kinetics of chylomicron metabolism from the metabolism of VLDLs in the absence of radiotracer and will not allow the isolation of naïve chylomicrons (since all the chylomicrons will be coated with poloxamer). Therefore, the mesenteric lymph is the most precise compartment for quantifying dietary lipid absorption and isolating chylomicrons.

Another advantage of isolating pure postlipid infusion mesenteric lymph is the ease with which chylomicrons can be isolated compared with from blood. We routinely ultracentrifuge lymph and simply collect the top creamy portion (53, 64, 94, 95). In postprandial lymph, this buoyant, density >1.006, fraction is almost all chylomicrons (63, 96). Mice secrete both apoB48 and apoB-100 on chylomicrons, whereas humans only make apoB-48 in the intestine, so we rarely use apoB-48 to differentiate murine lipoproteins (34). By comparison, isolating TAG-rich lipoproteins (chylomicrons and VLDL) from plasma is best done by tube slicing or differential centrifugation (69). In both cases, the chylomicron fraction will contain a heterogenous mixture of chylomicrons, chylomicron remnants, and VLDL.

We and others have a significant interest in intestinal immune cells, including those with the potential to travel between the gut and circulation. The lymphatic niche is also an essential barrier to the movement of enteric pathogens throughout the body. We undertook a basic analysis of T-cell populations and did not focus on specific effect/memory T-cell subsets. Our data show that those analyses would be possible in the future. That we find no differences in T-cell populations in prelipid and postlipid bolus lymph could be due to a lack of granularity, since we analyzed a single mixed sample made up of 6 h of postlipid bolus lymph (rather than individually analyzing each hour of lymph). Because of the reasonable volume of lymph collected each hour, these analyses would be possible in the future. Despite this, we would not expect huge differences in T-cell populations, since mesenteric lymph is cannulated after the lymph leaves the mLNs. Any immune cells that are trafficking between the lamina propria and the mLN or vice versa would not be present in this postnodal lymph, though naïve cells would be present. This supports the role of the mLN as the final mucosal firewall between activated immune cells or bacterial translocation and peripheral tissues (97–99). The mLN firewall is critical because escape of antigen-presenting cells from the gut into the circulation would stimulate naïve extraintestinal immune cells to respond to mucosal antigens (dietary, microbial, and self). The result would be highly inflammatory effector immune cells trafficking to the site of those antigens (the gut), which would result in autoimmune disorders like Crohn’s disease and ulcerative colitis. Our model is a unique biological site where appropriate versus

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**Fig. 5. (Continued).**
defective immune cell dynamics can be measured, and the postnodal lymph compartment could be a rich source of information about the inflammatory state of the mucosa in many diseases.

In addition to chylomicrons, the small intestine also secretes HDLs, fat soluble-vitamins, and antigen from commensal microorganisms. It is intriguing to speculate that comparisons of lymphocyte populations, specialized lipids, antigens, and gut hormones, in postnodal flowing lymph versus the blood compartment could reveal dynamics of intestinal immunity and mucosal barrier function, nutrient processing, and dysfunction in these processes during metabolic disease. We propose this 1-day lymph fistula model as a valuable tool for directly measuring these processes and show for the first time that these analyses are possible with the 1-day lymph fistula mouse model.

Data Availability
Raw data are housed on the Kohan Lab server at the University of Pittsburgh School of Medicine. Data will be shared upon request to the corresponding author: Alison B. Kohan, PhD, University of Pittsburgh School of Medicine; Email: akohan@pitt.edu.

Supplemental Data
This article contains supplemental data.

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Author Contributions
A. B. K. conceptualization; N. D. methodology; N. D. validation; L. T. and J. S. K. formal analysis; N. D. and L. T. investigation; N. D. resources; L. T. and J. S. K. data curation; A. B. K. writing—original draft; N. D. writing—review & editing; L. T. and J. S. K. visualization; A. B. K. supervision; A. B. K. project administration; A. B. K. funding acquisition.

Author ORCIDs
Alison B. Kohan https://orcid.org/0000-0003-3127-7283

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Conflict of Interest
The authors declare that they have no conflicts of interest with the contents of this article.

Abbreviations
FSC, forward scatter; FSC-A, forward scatter area; LPS, lipopolysaccharide; mLN, mesenteric lymph node; TAG, triglyceride.

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Supplemental Data
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