Abstract

Owing to the worldwide shortage of deceased-donor organs for transplantation, living donations have become a significant source of transplant organs. However, not all willing donors can donate to their intended recipients because of medical incompatibilities. These incompatibilities can be overcome by an exchange of donors between patients. For kidneys, such exchanges have become widespread in the last decade with the introduction of optimization and market design techniques to kidney exchange. A small but growing number of liver exchanges have also been conducted. Over the last two decades, a number of transplantation procedures emerged where organs from two living donors are transplanted to a single patient. Prominent examples include dual-graft liver transplantation, lobar lung transplantation, and simultaneous liver-kidney transplantation. Exchange, however, has been neither practiced nor introduced in this context. We introduce dual-donor organ exchange as a novel transplantation modality, and through simulations show that living-donor transplants can be significantly increased through such exchanges. We also provide a simple theoretical model for dual-donor organ exchange and introduce optimal exchange mechanisms under various logistical constraints.

Keywords: Market Design, Matching, Complementarities, Lung Exchange, Dual-Graft Liver Exchange, Simultaneous Liver-Kidney Exchange

JEL Codes: D47, C78
1 Introduction

Most transplants from living donors require only one donor for each procedure. There are, however, exceptions, including dual-graft liver transplantation, bilateral living-donor lobar lung transplantation, and simultaneous liver-kidney transplantation. For each of these procedures, grafts from two compatible living donors are transplanted. As such, these procedures are more involved from an organizational perspective than those with only one donor. Unfortunately, one or both of the donors can often be biologically incompatible with the intended recipient, precluding the transplantation. One way to overcome this potential barrier to transplantation is by an exchange of donors between patients. In addition to now-widespread kidney exchange, a small but growing number of (single-graft) liver exchanges has been conducted since the introduction of this transplantation modality in South Korea in 2003 (Hwang et al., 2010). Despite the introduction of two-donor transplantation techniques, living-donor organ exchange has not yet been practiced, or even introduced for these procedures. In this paper, we fill this gap as we

1. introduce dual-donor organ exchange as a potential transplantation modality for
   (a) dual-graft liver transplantation,
   (b) bilateral living-donor lobar lung transplantation, and
   (c) simultaneous liver-kidney transplantation,
2. simulate the gains from exchange based on data from South Korea (for the applications of dual-graft liver transplantation and simultaneous liver-kidney transplantation) and Japan (for the application of bilateral living-donor lobar lung transplantation),
3. develop a model of dual-donor organ exchange, and
4. introduce exchange mechanisms under various logistical constraints.

As in kidney exchange, all operations in dual-donor organ exchange have to be carried out simultaneously. This practice ensures that no donor donates an organ or a lobe unless his intended recipient receives a transplant. As such, organization of these exchanges is not an easy task: A 2-way exchange involves six simultaneous operations, a 3-way exchange involves nine simultaneous operations, and so on. As shown by Roth, Sönmez, and Ünver (2007), most of the gains from kidney exchange can be obtained by exchanges that are no larger than 3-way. In this paper, we show that this is not the case for dual-donor organ exchange. Our simulations suggest that number of transplants from larger than 3-way exchanges can approach to the number transplants from 2-way and 3-way exchanges combined (See Table 2). Therefore, exploring the structure of optimal exchange mechanisms is important under various constraints on the size of feasible exchanges.

---

1 Simulations are conducted for countries where the respective transplantation modality is most prominent.
Our model builds on the kidney-exchange model of Roth, Sönmez, and Ünver (2004, 2007). Medical literature\(^2\) suggests that a living donor can donate an organ or a lobe to a patient if he is

1. **blood-type compatible** with the patient for the cases of kidney transplantation, liver transplantation, and lung transplantation,

2. **size-compatible** (in the sense that the donor is at least as large as the patient) for the cases of single-graft liver transplantation and lobar lung transplantation, and

3. **tissue-type compatible** for the case of kidney transplantation.

For our simulations, we take all relevant compatibility requirements into consideration in order to assess the potential welfare gains from dual-donor organ exchange under various constraints. For our analytical results on optimal exchange mechanisms, we consider a simplified model with blood-type compatibility only. With this modeling choice, our analytical model captures all essential features of dual-graft liver transplantation and lobar lung transplantation for pediatric patients, but it is only an approximation for the applications of lobar lung transplantation for adult patients and simultaneous liver-kidney transplantation. Focusing on blood-type compatibility alone allows us to define each patient as a triple of blood types (one for the patient and two for her incompatible donors), making our model analytically tractable.

While there are important similarities between kidney exchange and dual-donor organ exchange, there are also major differences. From an analytical perspective, the most important difference is the presence of two donors for each patient rather than only one as in the case of kidney exchange. For each patient, the two donors are perfect complements.\(^3\) This key difference makes the dual-donor organ exchange model analytically more demanding than the (single-donor) kidney-exchange model. Even organizing an individual exchange becomes a richer problem under dual-donor organ exchange. For kidney exchange, each exchange (regardless of the size of the exchange) has a cycle configuration, where the donor of each patient donates a kidney to the next patient in the cycle. For dual-donor organ exchange, there are two configurations for a 2-way exchange (see Figure 1), five configurations for a 3-way exchange (see Figure 2), and so on. The richness of exchange configurations in our model also means that the optimal organization of these exchanges will be more challenging than for kidney exchange. Despite this technical challenge, we provide optimal mechanisms for (i) 2-way exchanges, (ii) 2-way and 3-way exchanges, and (iii) unrestricted exchanges (in Appendix E).

\(^2\)For commonly practiced compatibility requirements, see Lee et al. (2001) and Florman and Miller (2006) for liver transplantation and McLean, Barr, and Starnes (2007) and Van Raemdonck et al. (2009) for lung transplantation. The need for tissue-type compatibility for kidney transplantation is well established, while tissue-type compatibility is not required in general for liver transplantation (Cecka, Zhang, and Reed, 2005). There is no well-established protocol on tissue-type compatibility for lung transplantation.

\(^3\)In matching literature there are not many models that can incorporate complementarities and find positive results. Most of the matching literature focuses on various substitutability conditions and shows negative results even in the existence of slight complementarities in preferences. For example, see Hatfield and Milgrom (2005), Hatfield and Kojima (2008), and Hatfield and Kominers (2015).
Figure 1: Possible 2-way exchanges. Each patient (denoted by P) and her paired donors (each denoted by D) are represented in an ellipse. Carried donations in each exchange are represented by directed line segments. On the left, each patient swaps both of her donors with the other patient. On the right, each patient swaps a single donor with the other patient and receives a graft from her other donor.

Figure 2: Possible 3-way exchanges. On the upper-left, each patient trades one donor in a clockwise trade and the other donor in a counterclockwise trade. On the upper-right, each patient trades both of her donors in clockwise trades. On the lower-left, each patient trades one donor in a clockwise exchange and receives a graft from her other donor. On the middle, one patient is treated asymmetrically with respect to the other two: one patient trades both of her donors in two 2-way trades, one with one patient, the other with the other patient, while each of the other patients receives a graft from her remaining donor. On the lower-right, all patients are treated asymmetrically; one patient receives from one of her own donors, and one patient’s donors both donate to a single patient, while the last patient’s donors donate to the other two patients.

Due to compatibility requirements between a patient and each of her donors, living donation for dual-donor procedures proves to be a challenge to arrange even for patients with willing donors. But this friction also suggests that the role of an organized exchange can be more prominent for these procedures than for single-donor procedures. Our simulations in Section 3 and Appendix B
confirm this insight. An organized lung exchange in Japan has the potential to increase the number of living-donor lung transplants through 2-way and 3-way exchanges by 134%-200%, saving as many as 40 additional lung patients annually (see last line of Table 2 in Section 3.1 and Table 5 in Appendix B.1). Even though dual-graft liver transplantation is a secondary option to single-graft liver transplantation, an organized dual-graft liver exchange has a potential to increase the number of living-donor liver transplants by 23%-30% through 2-way and 3-way exchanges, saving nearly 230-300 additional liver patients in South Korea alone (see the last line of Table 4 in Section 3.2 and Table 6 in Appendix B.2).

Increasingly, economists are taking advantage of advances in technology to design new or improved allocation mechanisms in applications as diverse as entry-level labor markets (Roth and Peranson, 1999), spectrum auctions (Milgrom, 2000), internet auctions (Edelman, Ostrovsky, and Schwarz, 2007; Varian, 2007), school choice (Abdulkadiroğlu and Sönmez, 2003), kidney exchange (Roth, Sönmez, and Ünver, 2004, 2005, 2007), course allocation (Budish and Cantillon, 2012; Sönmez and Ünver, 2010), affirmative action (Echenique and Yenmez, 2015; Hafalir, Yenmez, and Yildirim, 2013; Kojima, 2012), cadet-branch matching (Sönmez, 2013; Sönmez and Switzer, 2013), refugee matching (Jones and Teytelboym, 2016; Moraga and Rapoport, 2014), and assignment of arrival slots (Abizada and Schummer, 2013; Schummer and Vohra, 2013). Our paper contributes to the emerging field of market design by introducing a new application in dual-donor organ exchange, and also to transplantation literature by introducing three novel transplantation modalities.

2 Background for Applications

There are four human blood types, $O$, $A$, $B$, and $AB$, denoting the existence or absence of the two blood proteins $A$ or $B$ in the human blood. A patient can receive a donor’s transplant organ (or a lobe of an organ), unless the donor carries a blood protein that the patient does not have. Thus, in the absence of other requirements, $O$ patients can receive a transplant from only $O$ donors, $A$ patients can receive a transplant from $A$ and $O$ donors, $B$ patients can receive a transplant from $B$ and $O$ donors, and $AB$ patients can receive a transplant from all donors. For some of our applications, there are additional medical requirements. In addition to the background information for each of these applications, the presence or lack of additional compatibility requirements are discussed below for dual-graft liver transplantation and living-donor lobar lung transplantation. Background for simultaneous liver-kidney transplantation is discussed in Appendix C.

2.1 Dual-Graft Liver Transplantation

The liver is the second most common organ for transplantation, after the kidney. Of nearly 31000 US transplants in 2015, more than 7000 were liver transplants. While there is the alternative (albeit inferior) treatment of dialysis for end-stage kidney disease, there are no alternatives to transplantation for end-stage liver disease. In contrast to western countries, donations for liver
transplantation in much of Asia come from living donors. For example in 2015, while only 359 of 7127 liver transplants were from living donors in the US, 942 of 1398 liver transplants were from living donors in South Korea. The low rates of deceased-donor organ donation in Asia are to a large extent due to cultural reasons and beliefs to respect bodily integrity after death (Lee, 2010). The need to resort to living-donor liver transplantation arose as a response to the critical shortage of deceased-donor organs and the increasing demand for liver transplantation in Asia, where the incidence of end-stage liver disease is very high (Lee et al., 2001). For similar reasons, living-donor liver transplantation is also more common than deceased-donor liver transplantation in several countries with predominantly Muslim populations, such as Turkey and Saudi Arabia.

A healthy human can donate part of his liver, which typically regenerates within a month. Donation of the smaller left lobe (normally 30-40% of the liver) or the larger right lobe (normally 60-70% of the liver) are the two main options. In order to provide adequate liver function for the patient, at least 40% and preferably 50% of the standard liver volume of the patient is required. The metabolic demands of a larger patient will not be met by the smaller left lobe from a relatively small donor. This phenomenon is referred to as small-for-size syndrome by the transplantation community. The primary solution to avoid this syndrome has been harvesting the larger right lobe of the liver. This procedure, however, is considerably more risky for the donor than harvesting the much smaller left lobe.\footnote{While donor mortality is approximately 0.1\% for left lobe donation, it ranges from 0.4\% to 0.5\% for right lobe donation (Lee, 2010). Other risks, referred to as donor morbidity, are also considerably higher with right lobe donation.} Furthermore, for donors with larger than normal-size right lobes, this option is not feasible.\footnote{For the donor, at least 30\% of the standard liver volume of the donor is required. Beyond this limit, the remnant liver of the donor loses its ability to compensate, regenerate, and recover (Lee et al., 2001).} Even though the patient receives an adequate graft volume with right lobe transplantation, the remaining left lobe may not be enough for donor safety. Thus, unlike deceased-donor whole-size liver transplantation, size matching between the liver graft and the standard liver volume of the patient has been a major challenge in adult living-donor liver transplantation due to the importance of providing an adequate graft mass to the patient while leaving a sufficient mass of remaining liver in the donor to ensure donor safety.

Dual-graft (or dual-lobe) liver transplantation, a technique that was introduced by Sung-Gyu Lee at the Asan Medical Center of South Korea in 2000, emerged as a response to the challenges of the more risky right lobe liver transplantation (Lee et al., 2001). Under this procedure, one (almost always left) liver lobe is removed from each of the two donors, and they are both transplanted into a patient. In the period 2011-2015, 176 dual-graft liver transplants were performed in South Korea, with the vast majority at the Asan Medical Center. Other countries that have performed dual-graft liver transplantation so far include Brazil, China, Germany, Hong Kong, India, Romania, and Turkey. The presence of two willing donors (almost always) solves the problem of size matching, rendering size-compatibility inconsequential, but transplantation cannot go through if one or both donors are blood-type incompatible with the patient. This is where an exchange of donors can play an important role, making dual-graft liver transplantation an ideal application for dual-donor
organ exchange with blood-type compatibility only. As an interesting side note, single-lobe liver exchange was introduced in 2003 at the Asan Medical Center, the same hospital where dual-graft liver transplantation was introduced. As such, it is a natural candidate to adopt an exchange program for potential dual-graft liver recipients.⁶

### 2.2 Living-Donor Lobar Lung Transplantation

As in the case of kidneys and livers, deceased-donor lung donations have not been able to meet the demand. As a result, thousands of patients worldwide die annually while waiting for lung transplantation. Living-donor lobar lung transplantation was introduced in 1990 by Dr. Vaughn Starnes and his colleagues for patients who are too critically ill to survive the waiting list for deceased-donor lungs. Since then, eligibility for this novel transplantation modality has been expanded to cystic fibrosis and other end-stage lung diseases.

A healthy human has five lung lobes: three lobes in the right lung and two in the left. In a living-donor lobar lung transplantation, two donors each donate a lower lobe to the patient to replace the patient’s dysfunctional lungs. Each donor must not only be blood-type compatible with the patient, but, donating only a part of the lung, he should also weigh at least as much. Hence blood-type compatibility and size compatibility are the two major medical requirements for living-donor lobar lung transplantation. This makes living donation much harder to arrange for lungs than for kidneys, even if a patient is able to find two willing donors.

Sato et al. (2014) report that there is no significant difference in patient survival between living-donor and deceased-donor lung transplantations. For a living donor, however, donation of part of a lung is “more costly” than donation of a kidney or even the left lobe of a liver. A healthy donor can maintain a normal life with only one kidney. And the liver regenerates itself within months after a living donation. In contrast, a donated lung lobe does not regenerate resulting in a loss of 10-20% of pre-donation lung capacity. In large part due to this discouraging reason, there have been only 15-30 living-donor lobar lung transplants annually in the US in the period 1994-2004. This already modest rate has essentially diminished in the US over the last decade as the lung allocation score (LAS) was initiated in May 2005 to allocate lungs on the basis of medical urgency and post-transplant survival. Prior to LAS, allocation of deceased-donor lungs was mostly based on a first-come-first-serve basis.

At present, Japan is the only country with a strong presence in living-donor lung transplantation. In 2013, there were 61 lung transplantations in Japan, of which 20 were from living donors. Okayama University hospital has the largest program in Japan, having conducted nearly half of the

---

⁶From an optimal design perspective, it would be preferable to combine our proposed dual-graft liver exchange program with the existing single-graft liver exchange program. When exchanges are restricted to logistically easier 2-way exchanges, such a unification can only be beneficial if a patient is allowed to receive a graft from a single donor in exchange for grafts from two of her donors. We leave this possibility to potential future research, in part because an exchange of “two donors for only one donor” has no medical precedence, and it may be subject to criticism by the medical ethics community.
living-donor lung transplants. Since September 2014, we have been collaborating with their lung-transplantation team to assess the potential of a lung-exchange program at Okayama University hospital.

3 Simulations

We start our analysis with calibrated simulations to quantify the potential gains from dual-donor organ exchange.\(^7\) Our methodology to generate patients and their attached donors is similar for all simulations. Each patient is randomly generated according to her respective population characteristics. For most applications, each patient is attached to two independently and randomly generated donors.\(^8\)

The construction of the dual-donor exchange pool depends on the specific application, the most straightforward one being the case of lung transplantation. For this application, any patient who is incompatible with one or both of her attached donors is sent to the exchange pool. Once the exchange pool forms, an optimal algorithm is used to determine the transplants via exchange to maximize the number of transplants. For liver transplantation, we assume that single-graft transplantation is preferred to dual-graft transplantation because the former puts only one donor in harm’s way rather than two. Therefore, for any patient, (i) direct donation from a single donor, (ii) exchange with a single donor, and (iii) direct donation from two donors will all be attempted in the given order before the patient is sent to the dual-graft liver-exchange pool.\(^9\) We also conduct dynamic simulations and report their findings in Appendix B.\(^10\)

3.1 Lung Exchange

Since Japan leads the world in living-donor lung transplantation, we simulate patient-donor characteristics based on data available from that country. We failed to obtain gender data for Japanese transplant patients. Therefore, we assumed that half of the patient population is male and the other half is female. We use the aggregate data statistics in Table 1 to calibrate the simulation parameters.\(^11\) Each patient-donor-donor triple is specified by their blood types and weights. We deem a patient compatible with a donor if the donor is blood-type compatible with the patient and

\(^{7}\)SLK-exchange simulations are reported in Appendix C.

\(^{8}\)For the case of simultaneous liver-kidney (SLK) exchange, there are kidney-only and liver-only patients who are in need of one donor only. A single donor is generated for these patients.

\(^{9}\)For the application of SLK transplantation, we consider a scenario where the exchange pool not only includes SLK patients who are incompatible with one or both of their donors, but also kidney (only) patients and liver (only) patients with incompatible donors.

\(^{10}\)See supplementary files for the Matlab program files and the data files used in simulations.

\(^{11}\)For random parameters like height, weight, or left-lobe liver volume percentage in liver transplantation, we only have the mean and standard deviation of the population distributions. Using these moments, we assume that these variables are distributed by a truncated normal distribution. The choices of the truncation points are \(\mu \pm c\sigma\) where \(\mu\) is the mean, \(\sigma\) is the standard deviation of the distribution, and coefficient \(c\) is set to 3 (a large number chosen not to affect the reported variance of the distributions much). The truncated normal distribution PDF with truncation
also as heavy as the patient. We consider population sizes of \( n = 10, 20, \) and \( 50 \) for the simulations.

| Lung Disease Patients 2013 |  |
|---------------------------|--|---|
| Waitlisted at the beginning of the year | Arrived/departed during the year | Received live-/deceased-donor trans. |
| 193 | 126–146 / 25–45 | 20 / 41 |

| Adult Body Weight (kg.) |  |
|-------------------------|--|---|
| Female | Mean: 52.9 | Std Dev: 9.0 |
| Male | Mean: 65.7 | Std Dev: 11.1 |
| Composite | Mean: 59.3 | Std Dev: 10.1 |

**Blood-Type Distribution**

<table>
<thead>
<tr>
<th>Type</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
<td>30.05%</td>
</tr>
<tr>
<td>A</td>
<td>40.00%</td>
</tr>
<tr>
<td>B</td>
<td>20.00%</td>
</tr>
<tr>
<td>AB</td>
<td>9.95%</td>
</tr>
</tbody>
</table>

**Table 1:** Summary statistics for lung-exchange simulations. This table reflects the parameters used in calibrating the simulations for lung exchange. We obtained the blood-type distribution for Japan from the Japanese Red Cross website [http://www.jrc.or.jp/donation/first/knowledge/index.html](http://www.jrc.or.jp/donation/first/knowledge/index.html) on 04/10/2016. The Japanese adult weight distribution’s mean and standard deviation were obtained from e-Stat of Japan using the 2010 National Health and Nutrition Examination Survey from the website [https://www.e-stat.go.jp/SG1/estat/GL02010101.do?method=init](https://www.e-stat.go.jp/SG1/estat/GL02010101.do?method=init) on 04/10/2016.

Patients who are compatible with both donors receive two lobes from their own donors directly, whereas the remaining patients join the exchange pool. Then we find optimal 2-way, 2&3-way, 2–4-way, 2–5-way, and unrestricted matchings.

Simulation results are reported in Table 2. When \( n = 50 \) (the last two lines), only 12.6% of the patients can receive direct donation, and the rest, 87.4%, participate in exchange (that is the remaining average of 43.7 patients). Using only 2-way exchange, an additional 10% of the patients can be matched, increasing the number of living-donor transplants by 78.5% (that is 4.96 divided by 6.31). Using 2&3-way exchanges, we can increase the number of living-donor transplants by 135% (that is 8.51 divided by 6.31). Of course, larger exchange sizes require more transplant teams to be simultaneously available and can test the limits of logistical constraints. Subject to this caveat, it is possible to match nearly 25% of all patients via 2–5-way exchanges, almost tripling the number of living-donor lung transplants. At the limit, i.e., in the absence of restrictions on exchange sizes, a third of the patients can receive lung transplants through exchanges, facilitating living-donor lung transplantation to nearly 46% of all patients in the population (matching 16.5 patients in exchange in addition to the 6.31 patients who receive direct transplantation).

The probability density function (PDF) and cumulative distribution function (CDF) of standard normal distribution, respectively, is given as

\[
 f(x; \mu, \sigma, a, b) = \frac{\frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{(x-\mu)^2}{2\sigma^2}}}{\Phi\left(\frac{b-\mu}{\sigma}\right) - \Phi\left(\frac{a-\mu}{\sigma}\right)},
\]

where \( \Phi \) and \( \Phi \) are the PDF and CDF of standard normal distribution, respectively.

9
The effect of the population size on marginal contribution of exchange is very significant: For example, the contribution of 2&3-way exchange to living-donor transplantation reduces from 135% to 30% when the population size reduces from $n = 50$ to $n = 10$.

<table>
<thead>
<tr>
<th>Population Size</th>
<th>Direct Donation</th>
<th>Exchange Technology</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2-way</td>
<td>2&amp;3-way</td>
</tr>
<tr>
<td>10</td>
<td>1.256</td>
<td>0.292</td>
</tr>
<tr>
<td></td>
<td>(1.0298)</td>
<td>(0.72925)</td>
</tr>
<tr>
<td>20</td>
<td>2.474</td>
<td>1.128</td>
</tr>
<tr>
<td></td>
<td>(1.4919)</td>
<td>(1.4183)</td>
</tr>
<tr>
<td></td>
<td>(2.962)</td>
<td>(2.9759)</td>
</tr>
</tbody>
</table>

Table 2: Lung-exchange simulations. In these results and others, the sample standard deviations reported are reported under averages; for the standard errors of the averages, these deviations need to be divided by the square root of the simulation number, $\sqrt{500} = 22.361$.

### 3.2 Dual-Graft Liver Exchange

For simulations on dual-graft liver exchange, we use the South Korean population characteristics (see Table 3). The same statistics are used for the SLK-exchange simulations in Appendix C.

We restrict our attention to left-lobe transplantation only, a procedure that is considerably safer for the donor than right-lobe transplantation. The Korean adult liver left lobe volume distribution’s moments are also given in Um et al. (2015) (see Table 3). We randomly set the graft volume of each donor using these parameters. We consider the following simulation scenario in given order, as dual-graft liver transplants are considered only if a suitable single-graft donor cannot be found:

1. If at least one of the donors of the patient is blood-type compatible, and his graft volume is at least 40% of the liver volume of the patient, then the patient receives a transplant directly from this compatible donor (denoted as “1-donor direct” scenario).

---

12 There is a selection bias in using the gender distributions of who receive transplants and who donate, instead of those of who need transplants and who volunteer to donate. We use the former in our simulations because this is the best data publicly available and we did not want to speculate about the underlying gender specific disease and behavioral donation models that generate the latter statistics.

13 In generating patient populations, we assume that each patient is attached to two living donors. We determine the blood type, gender, and height characteristics for patients and their donors independently and randomly. Then, we use the following weight determination formula as a function of height: $w = ah^b$, where $w$ is weight in kg, $h$ is height in meters, and constants $a$ and $b$ are set as $a = 26.58, b = 1.92$ for males and $a = 32.79, b = 1.45$ for females (Diverse Populations Collaborative Group, 2005). This is the best formula we could find for a weight-height relationship in humans in this respect. This paper does not report confidence intervals; therefore, we could not use a stochastic process to generate weights. The body surface area ($BSA$ in m$^2$) of an individual is determined through the Mostellar formula given in Um et al. (2015) as $BSA = \sqrt{\frac{hw}{6}}$, and the liver volume ($l_v$ in ml) of Korean adults is determined through the estimated formula in Um et al. (2015) as $l_v = 893.485 BSA - 439.169$. 

10
<table>
<thead>
<tr>
<th>Statistics for Dual-Graft Liver-Exchange from South Korean Population</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Live-Liver Donation Recipients in 2010-2014</strong></td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Total</td>
</tr>
<tr>
<td><strong>Live-Liver Donors in 2010-2014</strong></td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Adult Height (cm.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female Mean: 157.4 Std Dev: 5.99</td>
</tr>
<tr>
<td>Male Mean: 170.7 Std Dev: 6.4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Liver Left Lobe Volume as Percentage of Whole</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean: 34.7% Std Dev: 3.9%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Blood-Type Distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>O</td>
</tr>
<tr>
<td>A</td>
</tr>
<tr>
<td>B</td>
</tr>
<tr>
<td>AB</td>
</tr>
</tbody>
</table>

Table 3: Summary statistics for dual-graft liver-exchange simulations. This table reflects the parameters used in calibrating the simulations. We obtained the blood-type distribution for South Korea from [http://bloodtypes.jigsy.com/East_Area-bloodytypes](http://bloodtypes.jigsy.com/East_Area-bloodytypes) on 04/10/2016. The South Korean adult height distribution’s mean and standard deviation were obtained from the Korean Agency for Technology and Standards (KATS) website [http://sizekorea.kats.go.kr](http://sizekorea.kats.go.kr) on 04/10/2016. The transplant data were obtained from the Korean Network for Organ Sharing (KONOS) 2014 Annual Report, retrieved from [https://www.konos.go.kr/konosis/index.jsp](https://www.konos.go.kr/konosis/index.jsp) on 04/10/2016.

2. The remaining patients and their donors participate in an optimal “1-donor exchange” program. We use the same criterion as above to determine compatibility between any patient and any donor in the 1-donor exchange pool. Specifically, patients form 2-way (or 2&3 way) exchanges in which each patient receives a graft that is at least 40% of her liver volume from a blood-type compatible donor of another patient in the same exchange.

3. The remaining patients and their coupled donors are checked for dual-graft compatibility. If a patient’s donors are blood-type compatible with her and the sum of the donors’ graft volumes is at least 40% of the patient’s liver volume, then the patient receives dual grafts from her own donors (denoted as “2-donor direct” scenario).

4. Finally the remaining patients and their donors participate in an optimal “2-donor exchange” program. We use the same criterion as above to deem any pair of donors dual-graft compatible with any patient. Specifically, patients form 2-way (or 2&3 way) exchanges in which each patient receives two grafts that total to at least 40% of her liver volume from two blood-type compatible donors, at least one of whom is paired with a different patient in the same exchange.
Simulation results are reported in Table 4. For a population of $n = 250$ (in the last 6 lines in the table), on average 141 patients remain without a transplant following 1-donor direct transplant and 1-donor 2&3-way exchange modalities (as about 60 patients receive transplant from a donor of theirs and an additional 49 patients receive 1-donor exchange transplants as seen in the third line of $n = 250$). About 31% of these patients receive dual-graft transplants from their own donors under the 2-donor direct modality (that is around 43.5 patients receive 2-donor transplants from their own donors out of the 141 remaining). An additional 24.5% of these patients are matched in the 2-donor 2&3-way exchange modality (that is around 35 additional patients receive 2-donor transplants through exchange out of the 141 remaining). This final figure corresponds to approximately 80% of the 2-donor direct donation, and thus, the contribution of exchange to dual-graft transplantation is highly significant. Moreover, the 2-donor 2&3-way exchange modality provides transplants for 70.5% of the number of patients who receive transplants through the 1-donor exchange modality. Therefore, the contribution of the 2-donor exchange modality to the overall number of transplants from exchange is also highly significant. Under 2&3-way exchanges, 2-donor exchange increases the total number of living-donor liver transplants by about 23% by matching 13.9% of all patients.

Since pools evolve differently depending on which exchange size constraint is used, we also include three columns titled “As % of Entrants” in Table 4, which report patients who receive transplants under each modality (1-Donor Exchange, 2-Donor Direct, or 2-Donor Exchange) as the percentage of the patients who are present in the pool for the given transplant modality. Thus, the percentages in the last column can be used to compare gains from 2-Donor exchange for different population sizes and exchange-size constraints. For $n = 250$, by 2-way exchanges only, about 24.5% of the patients entering 2-Donor exchange receive transplants (i.e., 26 divided by 106, which is $n = 250$ minus the sum of patients matched in previous stages). By 2&3-way exchanges 35.75% of the patients entering 2-Donor exchange pool receive transplants, and in the absence of exchange-size constraints 38.7% of the patients entering 2-Donor exchange pool receive transplants. Thus, unlike in lung exchange, most of the gains from exchange are captured through 2&3-way exchanges in dual-graft liver exchange.

4 A Model of Dual-Donor Organ Exchange

Our simulations in Section 3 show that exchange is potentially important in the context of dual-donor organ transplants. We next present a simple theoretical model for its analysis.

We assume that each patient, who has two living donors, can receive transplant organs from her own donors if and only if both of them are blood-type compatible with the patient. That is, the two transplant organs are perfect complements for the patient. In our benchmark model, we assume that there are no size or tissue-type compatibility requirements; the only compatibility requirement regards the blood type. This assumption helps us to focus exclusively on the effect of the two-donor
Table 4: Dual-graft liver-exchange simulations.

<table>
<thead>
<tr>
<th>Population Size</th>
<th>1-Donor Exchange</th>
<th>2-Donor Exchange</th>
<th>2-Donor Exchange</th>
</tr>
</thead>
<tbody>
<tr>
<td>Direct</td>
<td>Matched</td>
<td>As % of Entrants</td>
<td>Matched</td>
</tr>
<tr>
<td>50</td>
<td>2-way</td>
<td>3.92 (2.7204)</td>
<td>10.33%</td>
</tr>
<tr>
<td></td>
<td>2&amp;3-way</td>
<td>5.066 (3.4382)</td>
<td>13.35%</td>
</tr>
<tr>
<td></td>
<td>Unrestricted</td>
<td>5.772 (3.9799)</td>
<td>15.21%</td>
</tr>
<tr>
<td>100</td>
<td>2-way</td>
<td>10.656 (4.2073)</td>
<td>14.04%</td>
</tr>
<tr>
<td></td>
<td>2&amp;3-way</td>
<td>14.452 (5.6152)</td>
<td>19.04%</td>
</tr>
<tr>
<td></td>
<td>Unrestricted</td>
<td>17.754 (6.4827)</td>
<td>23.39%</td>
</tr>
<tr>
<td>250</td>
<td>2-way</td>
<td>35.032 (7.5297)</td>
<td>18.44%</td>
</tr>
<tr>
<td></td>
<td>2&amp;3-way</td>
<td>49.198 (10.37)</td>
<td>25.89%</td>
</tr>
<tr>
<td></td>
<td>Unrestricted</td>
<td>60.672 (11.127)</td>
<td>31.93%</td>
</tr>
</tbody>
</table>

requirement on organ exchange, and it best fits our application of dual-graft liver transplantation.\textsuperscript{14}

Let $B = \{O, A, B, AB\}$ be the set of blood types. We denote generic elements by $X, Y, Z \in B$. Let $\succeq$ be the partial order on blood types defined by $X \succeq Y$ if and only blood type $X$ can donate to blood type $Y$. Figure 3 illustrates the partial order $\succeq$.\textsuperscript{15}

\begin{figure}
\centering
\begin{tikzpicture}
  \node (O) at (0,0) {O};
  \node (A) at (-1,1) {A};
  \node (B) at (1,1) {B};
  \node (AB) at (0,-1) {AB};
  \draw (O) -- (A);
  \draw (O) -- (B);
  \draw (A) -- (AB);
  \draw (B) -- (AB);
\end{tikzpicture}
\caption{The Partial Order $\succeq$ on the Set of Blood Types $B = \{O, A, B, AB\}$.}
\end{figure}

\textsuperscript{14}When first introduced, the target population for lobar lung transplantation was pediatric patients. Since the lung graft needs of children are not as voluminous as those of adults, the application of lobar lung transplantation also fits our model well when the exchange pool consists of pediatric patients.

\textsuperscript{15}For any $X, Y \in B$, $X \succeq Y$ if and only if there is a downward path from blood type $X$ to blood type $Y$ in Figure 3.
Each patient participates in the exchange with two donors, which we refer to as a **triple**. The relevant information concerning the patient and her two donors can be summarized as a triple of blood types $X - Y - Z \in B^3$, where $X$ is the blood type of the patient, and $Y$ and $Z$ are the blood types of the donors. We will refer to each element in $B^3$ as a **triple type** such that the order of the donors has no relevance. For example, an $O$ patient with a pair of $A$ and $B$ donors counts as both a triple of type $O - A - B$ and also a triple of type $O - B - A$.

**Definition 1** An exchange pool is a vector of nonnegative integers $E = \{n(X - Y - Z) : X - Y - Z \in B^3\}$ such that:

1. $n(X - Y - Z) = n(X - Z - Y)$ for all $X - Y - Z \in B^3$.
2. $n(X - Y - Z) = 0$ for all $X - Y - Z \in B^3$ such that $Y \succeq X$ and $Z \succeq X$.

The number $n(X - Y - Z)$ stands for the number of participating $X - Y - Z$ triples.

The first condition in the definition of an exchange pool corresponds to the assumption that the order of the donors does not matter, i.e., $X - Y - Z$ and $X - Z - Y$ represent the same type. The second condition corresponds to the assumption that compatible patient-donor triples do not participate in the exchange.

## 5 2-way Exchange

In this section, we assume that only 2-way exchanges are allowed. We characterize the maximum number of patients receiving transplants for any given exchange pool $E$. We also describe an algorithm that achieves this maximum.

A 2-way exchange is the simplest form of dual-donor organ exchange, involving two triples exchanging one or both of their donors’ grafts, and it is the easiest to coordinate. Thus, as a first step in our analysis, it is important to understand the structure and size of optimal matchings with only 2-way exchanges. There are forty types of triples after accounting for repetitions due to the reordering of donors. The following Lemma simplifies the problem substantially by showing that only six of these types may take part in 2-way exchanges. All proofs are relegated to Appendices A, D, and E.

**Lemma 1** In any given exchange pool $E$, the only types that could be part of a 2-way exchange are $A - Y - B$ and $B - Y - A$ where $Y \in \{O, A, B\}$.

---

16It is straightforward to integrate into our model patients who have one donor and who need one organ. We can do so by treating these patients as part of a triple where a virtual donor is of the same blood type as the patient.

17While only six of forty types can participate in 2-way exchanges, nearly half of the patient populations in our simulations belong to these types due to very high rates of blood types A and B in Japan and South Korea.
The six types of triples in Lemma 1 are such that every $A$ blood-type patient has at least one $B$ blood-type donor, and every $B$ blood-type patient has at least one $A$ blood-type donor. Therefore, $A$ blood-type patients can only take part in a 2-way exchange with $B$ blood-type patients, and vice versa. Furthermore, if they participate in a 2-way exchange, the $A-A-B$ and $B-B-A$ types must exchange exactly one donor; the $A-B-B$ and $B-A-A$ types must exchange both donors; and the $A-O-B$ and $B-O-A$ types might exchange one or two donors.

We refer the six types in Lemma 1 as **essential types** and summarize the possible 2-way exchanges between them as the edges of the graph in Figure 4.

We next present a matching algorithm that maximizes the number of transplants through 2-way exchanges. The algorithm sequentially maximizes three subsets of 2-way exchanges:

**Algorithm 1 (Sequential Matching Algorithm for 2-way Exchanges)**

**Step 1:** Match the maximum number of $A - A - B$ and $B - B - A$ types. Match the maximum number of $A - B - B$ and $B - A - A$ types.

**Step 2:** Match the maximum number of $A - O - B$ types with any subset of the remaining $B - B - A$ and $B - A - A$ types. Match the maximum number of $B - O - A$ types with any subset of the remaining $A - A - B$ and $A - B - B$ types.

**Step 3:** Match the maximum number of the remaining $A - O - B$ and $B - O - A$ types.

Figure 5 graphically illustrates the pairwise exchanges that are carried out at each step of the sequential matching algorithm. The mechanics of this algorithm is very intuitive and based on optimizing the flexibility offered by blood-type $O$ donors. Initially, the optimal use of triples endowed with blood-type $O$ donors is not clear, and for Step 1 they are “put on hold.” In this first step, as many triples as possible are matched without using any triple endowed with a blood-type $O$ donor. By Step 2, the optimal use of triples endowed with blood-type $O$ donors is revealed. In this step, as many triples as possible are matched with each other by using only one blood-type $O$
The next Theorem shows the optimality of Algorithm 1 and characterizes the maximum number of transplants through 2-way exchanges.

**Theorem 1**  Given an exchange pool $E$, Algorithm 1 maximizes the number of 2-way exchanges. The maximum number of patients receiving transplants through 2-way exchanges is 
$$2 \min\{N_1, N_2, N_3, N_4\}$$

where:

$$N_1 = n(A - A - B) + n(A - O - B) + n(A - B - B)$$

$$N_2 = n(A - O - B) + n(A - B - B) + n(B - B - A) + n(B - O - A)$$

$$N_3 = n(A - A - B) + n(A - O - B) + n(B - O - A) + n(B - A - A)$$

$$N_4 = n(B - B - A) + n(B - O - A) + n(B - A - A)$$

Figure 6 depicts the sets of triple types whose market populations are $N_1$, $N_2$, $N_3$, and $N_4$.

**Figure 6:** The Optimal 2-way Sequential Matching Algorithm

**Figure 6:** Potential Bottleneck Groups for Maximum Number of 2-way Exchanges

### 6 Larger-Size Exchanges

We have seen that when only 2-way exchanges are allowed, every 2-way exchange must involve exactly one $A$ and one $B$ blood-type patient. The following Lemma generalizes this observation to
$K$-way exchanges for arbitrary $K \geq 2$. In particular, every $K$-way exchange must involve an $A$ and a $B$ blood-type patient, but if $K \geq 3$, then it might also involve $O$ blood-type patients.

**Lemma 2** Fix $E$ and let $K \geq 2$. Then, the only types that could be part of a $K$-way exchange are $O - Y - A$, $O - Y - B$, $A - Y - B$, and $B - Y - A$ where $Y \in \{O, A, B\}$. Furthermore, every $K$-way exchange must involve an $A$ and a $B$ blood-type patient.

In kidney-exchange pools, $O$ patients with $A$ donors are much more numerous than their opposite type pairs, $A$ patients with $O$ donors. That is because $O$ patients with $A$ donors arrive for exchange all the time, while $A$ patients with $O$ donors only arrive if there is tissue-type incompatibility between them (as otherwise the donor is compatible and donates directly to the patient). This empirical observation is caused by the blood-type compatibility structure. In general, patients with less-sought-after blood-type donors relative to their own blood type become in excess and plentiful as the exchange pool grows in size. A similar situation will also occur in dual-donor organ exchange pools. For kidney-exchange models, Roth, Sönmez, and Ünver (2007) make an explicit long-run assumption regarding this asymmetry. We will make a corresponding assumption for dual-donor organ exchange below. However, our assumption will be milder as it will be imposed only for two types of triples rather than all triple types with less-sought-after donor blood types than their patients.

**Definition 2** An exchange pool $E$ satisfies the long-run assumption if for every feasible matching in the absence of exchange-size restrictions, there is at least one $O - O - A$ and one $O - O - B$ type that do not take part in any exchange.

Suppose that the exchange pool $E$ satisfies the long-run assumption and $\mu$ is a matching composed of any size exchanges. The long-run assumption ensures that we can create a new matching $\mu'$ from $\mu$ by replacing every $O - A - A$ or $O - A - B$ type taking part in an exchange with an unmatched $O - O - A$ type, and every $O - B - B$ type taking part in an exchange by an unmatched $O - O - B$ type. Then, the new matching $\mu'$ is composed of the same size exchanges as $\mu$, and it induces the same number of transplants as $\mu$. Furthermore, the only $O$ blood-type patients matched under $\mu'$ belong to the triples of types $O - O - A$ or $O - O - B$.

Let $\bar{K} \geq 2$ be the maximum allowed exchange size. Consider the problem of finding an optimal matching, i.e., one that maximizes the number of transplants when only $2, \ldots, \bar{K}$-way exchanges are feasible. By the above paragraph, for any optimal matching $\mu$, we can construct another optimal matching $\mu'$ in which the only triples with $O$ blood-type patients matched under $\mu'$ are of types $O - O - A$ or $O - O - B$. We summarize this observation as the following Lemma:

**Lemma 3** Let $\bar{K} \geq 2$ be the maximum allowed exchange size and let the exchange pool $E$ satisfy the long-run assumption. Then, there exists an optimal matching exclusively involving the two types $O - O - A$, $O - O - B$ and the six essential types $A - Y - B$, $B - Y - A$ where $Y \in \{O, A, B\}$. 

17
Also observe that, since the numbers of type $O-O-A$ and type $O-O-B$ triples are nonbinding by the long-run assumption, an optimal matching can be characterized just in terms of the numbers of the six essential types. In the next subsection we use this approach to describe an algorithm that achieves the maximum number of transplants when $\bar{K} = 3$.

### 6.1 2- & 3-way Exchanges

We continue our analysis with a characterization of the types that can be part of a 3-way exchange. It turns out that ruling out the types $O-A-A$ and $O-B-B$ in the construction of an optimal matching is without loss of generality for the case of 3-way exchange. Not only these triples cannot be matched under an optimal matching, they cannot be part of any 3-way exchange.

**Lemma 4** Given an exchange pool $E$, triples of types $O-A-A$ or $O-B-B$ cannot participate in any 3-way exchange.

Thus, the only types that can participate in a 3-way exchange are $O-O-A$, $O-O-B$, $O-A-B$, and the six essential types $A-Y-B$, $B-Y-A$ where $Y \in \{O, A, B\}$.

For expositional simplicity, next we describe a collection of 2- and 3-way exchanges divided into three groups. We show in Lemma 6 in Appendix A that one can restrict attention to these exchanges when constructing an optimal matching.

![Three Groups of 2- and 3-way Exchanges](image)

**Figure 7:** Three Groups of 2- and 3-way Exchanges

**Definition 3** Given an exchange pool $E$, a matching is in **simplified form** if it consists of exchanges in the following three groups:

---

19Without additional structure, finding an optimal matching for fixed $\bar{K} \geq 3$ is a computationally hard problem in the sense that it is NP-complete (Abraham, Blum, and Sandholm, 2007). In contrast, exploiting the structure imposed above, we will provide a polynomial-time algorithm for $\bar{K} = 3$ in the next subsection and for $\bar{K} = 6$ in Appendix E.
Group 1: 2-way exchanges exclusively involving types $A - A - B$, $A - B - B$, $B - B - A$, and $B - B - A$. These exchanges are represented in Figure 7 by a regular (i.e. nonbold/nondotted end) edge between two of these types.

Group 2: 3-way exchanges exclusively involving types $A - A - B$, $A - B - B$, $B - B - A$, and $B - B - A$ represented in Figure 7 by an edge with one dotted and one nondotted end. A 3-way exchange in this group consists of two triples of the type at the dotted end and one triple of the type at the nondotted end.

Group 3: 3-way exchanges involving two of the types $A - A - B$, $A - B - B$, $B - B - A$, $B - O - A$, $B - A - A$, and one of the types $O - O - A$, $O - O - B$, $O - A - B$. These exchanges are represented in Figure 7 by a bold edge between the former two types. 

We will show that when the long-run assumption is satisfied, the following matching algorithm maximizes the number of transplants through 2- and 3-way exchanges. The algorithm sequentially maximizes three subsets of exchanges:

Algorithm 2 (Sequential Matching Algorithm for 2- and 3-way Exchanges)

Step 1: Carry out group 1, group 2 exchanges in Figure 7 among types $A - A - B$, $A - B - B$, $B - B - A$, and $B - A - A$ to maximize the number of transplants subject to the following constraints (*):

1. Leave at least a total $\min\{n(A - A - B) + n(A - B - B), n(B - O - A)\}$ of $A - A - B$ and $A - B - B$ types unmatched.
2. Leave at least a total $\min\{n(B - B - A) + n(B - A - A), n(A - O - B)\}$ of $B - B - A$ and $B - A - A$ types unmatched.

Step 2: Carry out the maximum number of 3-way exchanges in Figure 7 involving $A - O - B$ types and the remaining $B - B - A$ or $B - A - A$ types. Similarly carry out the maximum number of 3-way exchanges involving $B - O - A$ types and the remaining $A - A - B$ or $A - B - B$ types.

Step 3: Carry out the maximum number of 3-way exchanges in Figure 7 involving the remaining $A - O - B$ and $B - O - A$ types.

Figure 8 graphically illustrates the 2- and 3-way exchanges that are carried out at each step of the sequential matching algorithm. The intuition for our second algorithm is slightly more involved.

---

20 There are five exchanges represented by bold edges in Figure 7. Four of these exchanges involve only one triple with an $O$ donor. For those exchanges, the third triple of the 3-way exchange is uniquely defined either of type $O - O - A$ or of type $O - O - B$. The fifth exchange represented by a bold edge in Figure 7 has two triples with an $O$ donor each. For this exchange, the third triple can be of any of the types $O - O - A$, $O - O - B$, or $O - A - B$.

21 For each of these 3-way exchanges, the third triple is uniquely defined either of type $O - O - A$ or of type $O - O - B$.

22 For these exchanges, the third triple could be of any of the types $O - O - A$, $O - O - B$, or $O - A - B$. 

Figure 8: The Optimal 2- and 3-way Sequential Matching Algorithm

When only 2-way exchanges are allowed, the only perk of a blood-type $O$ donor is in his flexibility to provide a transplant organ to either an $A$ or a $B$ patient. When 3-way exchanges are also allowed, a blood-type $O$ donor has an additional perk: He can help save an additional patient of blood type $O$. For example, a triple of type $A - O - B$ can be paired with a triple of type $B - B - A$ to save one additional triple of type $O - O - A$. Since each patient of type $A - O - B$ is in need if a patient of either type $B - B - A$ or type $B - A - A$ to save an extra patient through the 3-way exchange, the maximization in Step 1 has to be constrained. Otherwise a 3-way exchange would be sacrificed for a 2-way exchange, reducing the number of transplants. The rest of the mechanics is similar between the two algorithms. For expositional purposes, we present the subalgorithm that solves the constrained optimization in Step 1 in Appendix D. The following Theorem shows the optimality of Algorithm 2.

**Theorem 2** Given an exchange pool $\mathcal{E}$ satisfying the long-run assumption, Algorithm 2 maximizes the number of transplants through 2- and 3-way exchanges.

### 6.2 Necessity and Sufficiency of 6-way Exchanges

Although larger exchanges are logistically harder to organize, it is of theoretical interest to understand their potential role in dual-donor organ exchange. We next identify the types that can participate in an optimal matching in the absence of constraints on exchange size. In this setting, the only blood-type $O$ patients that can be part of an optimal matching are of types $O - O - A$ or $O - O - B$.

**Lemma 5** Given an exchange pool $\mathcal{E}$ satisfying the long-run assumption, $O - A - A$, $O - A - B$, and $O - B - B$ type triples are never matched in an optimal matching in the the absence of exchange-size constraints.

Thus, the only types that can participate in an optimal exchange are $O - O - A$, $O - O - B$, and the six essential types $A - Y - B$, $B - Y - A$ where $Y \in \{O, A, B\}$.
In our next result, we show that restricting attention to 2–6-way exchanges is sufficient to maximize the number of transplants through exchange. As part of the proof of this result, given in Appendix E, we also provide an algorithm that achieves the maximum.

**Theorem 3** Suppose that the exchange pool $\mathcal{E}$ satisfies the long-run assumption and exchange sizes are unrestricted. Then there exists an optimal matching that consists of exchanges no larger than 6-way.

The following example shows that using 6-way exchanges is not only sufficient, but also necessary to find an optimal matching for some exchange pools.

**Example 1** Consider an exchange pool with one triple of type $A - O - B$, two triples of type $B - O - A$, and three triples of $O - O - B$.\(^{23}\) Observe that all patients can receive two transplant organs of their blood type. Therefore, all patients are matched under an optimal matching. With three blood-type $O$ patients and six blood-type $O$ donors, all blood-type $O$ organs must be transplanted to blood-type $O$ patients (for otherwise not each patients would receive a transplant). This in turn implies that the two blood-type $A$ organs must be transplanted to the only blood-type $A$ patient. Hence, the triple of type $A - O - B$ should be in the same exchange as the two triples of type $B - O - A$. Equivalently, all triples with a non-$O$ patient should be part of the same exchange. But patients of $O - O - B$ triples each are in need of an additional blood-type $O$ donor, and thus $O - O - B$ triples should also be part of the same exchange. Hence, 6-way exchange is necessary to match all patients and obtain an optimal matching (see Figure 9).

\[\text{Figure 9: An optimal matching for the pool in Example 1 always consists of a single 6-way exchange.}\]

Our simulations in Section 3 suggest that while the total number of transplants from exchanges larger than 3-way can be significant and approach those from 2-way and 3-way exchanges combined

\(^{23}\)For the example to use the long run assumption, we can assume there are also one additional $O - O - B$ triple and an $O - O - A$ triple. In this case, at least one $O - O - B$ and this $O - O - A$ triple remains unmatched in every matching.
for the application of lung exchange, they are relatively modest for the application of dual-graft liver exchange. The contrast between the two sets of simulations suggests that the presence of size-compatibility increases the role of larger than 3-way exchanges. Our theoretical results in the absence of size compatibility (and their proofs) provide some insight for the relatively modest role of larger than 3-way exchanges for the simulations on dual-graft liver exchange. Proposition 2 in Appendix E shows that when there are no exchange-size constraints, an optimal matching can be constructed using only 2- and 3-way exchanges provided that there are no blood-type O patients in the pool. Moreover, by Lemma 5, triples of type $O - O - A$ and $O - O - B$ are the only triples with blood type $O$ patients, who can be matched in an optimal matching. Each triple of these types requires a second $O$ donor, which can only be supplied by $A - O - B$ or $B - O - A$ types. Thus, all welfare gains from exchanges beyond 3-way come from the ability of matching additional $O - O - A$ and $O - O - B$ types through the utilization of $A - O - B$ and $B - O - A$ types. In the absence of exchange-size constraints, a triple of $O - O - A$ or $O - O - B$ can be appended to any exchange for each triple of $A - O - B$ or $B - O - A$ that is part of the exchange (see for example Figure 9 where three types of $O - O - B$ are appended to three types of $A - O - B$ or $B - O - A$). When only 2- and 3-way exchanges are feasible, only one triple of types $O - O - A$ or $O - O - B$ can be appended to any 2-way exchange that includes one or two triples of types $A - O - B$ and $B - O - A$. But that means, larger than 3-way exchanges can only increase the total number of transplants by the difference between (1) the maximum number of $A - O - B$ and $B - O - A$ types that can be matched in the absence of exchange-size constraints and (2) the maximum number of distinct 2-way exchanges that includes $A - O - B$ or $B - O - A$ types when no exchange can be larger than 3-way. The two types $A - O - B$ and $B - O - A$ are essential and they already play a key role under 2- and 3-way exchange. Hence the difference cannot be very high, limiting the role of larger than 3-way exchanges in our theoretical model as well as in our application of dual-graft liver exchange.

7 Conclusion

For any organ with the possibility of living-donor transplantation, living-donor organ exchange is also medically feasible. Despite the introduction and practice of transplant procedures that require two donors, organ exchange in this context is neither discussed in the literature nor implemented in practice. We propose dual-donor organ exchange as a new transplantation modality, focusing on the following three transplantation procedures: dual-graft liver transplantation, bilateral living-donor lobar lung transplantations, and simultaneous liver-kidney transplantation. We simulate the potential gains from dual-donor exchanges for these applications. We also formulate an analytical model of dual-donor organ exchange and introduce optimal exchange mechanisms under various logistical constraints.

Analytically, dual-donor organ exchange is a more challenging problem than kidney exchange since each patient is in need of two compatible donors who are perfect complements. Exploiting
the structure induced by the blood-type compatibility requirement for organ transplantation, we introduce optimal exchange mechanisms under various logistical constraints. Abstracting away from additional medical compatibility considerations such as size compatibility and tissue-type compatibility, our analytical model best captures the specifics of dual-graft liver transplantation. For our calibrated simulations, however, we take into account these additional compatibility requirements (whenever relevant) for each application. Through these simulations we show that the marginal contribution of exchange to living-donor organ transplantation is very substantial. For example, adopting 2-way exchanges alone has the potential to increase the number of living-donor lung transplants by 78.5% in Japan (see Table 2).

The potential of an organized exchange for each medical application in a given society will likely depend on the following factors:

1. Availability and expertise in the required transplantation technique.
2. Prominence of living donation.
3. Legal and cultural attitudes towards living-donor organ exchanges.

First and foremost, transplantation procedures that require two living donors are highly specialized and so far they are available only in a few countries. For example, the practice of living-donor lobar lung transplantation is reported in the literature only in the US and Japan. Hence, the availability of the required transplantation technology limits the potential markets for applications of dual-donor organ exchange. Next, organized exchange is more likely to succeed in an environment where living-donor organ transplantation is the norm rather than an exception. While living donation of kidneys is widespread in several western countries, it is much less common for organs that require more invasive surgeries, such as the liver and the lung. Since all our applications rely on these more invasive procedures, this second factor further limits the potential of organized exchange in the western world. In contrast, this factor is very favorable in several Asian counties and countries with predominantly Muslim populations, where living donors are the primary source of transplant organs. Finally, the concept of living-donor organ exchange is not equally accepted throughout the world, and it is not even legal in some countries. For example, organ exchanges are outlawed under the German transplant law. Indeed, it was unclear whether kidney exchanges violate the National Organ Transplant Act of 1984 in the US until Congress passed the Charlie W. Norwood Living Organ Donation Act of 2007, clarifying them as legal. Clearly dual-donor organ exchanges cannot flourish in a country unless they comply with the laws.

Based on these factors, we foresee the strongest potential for organized exchange for dual-graft liver transplantation in South Korea, for lobar lung transplantation in Japan, and for simultaneous liver-kidney transplantation in South Korea and in Turkey.
Appendix A  Proofs of Lemmas 1, 2, 4, & 5 and Theorems 1 & 2

Proof of Lemma 1: Since $AB$ blood-type patients are compatible with their donors, there are no $AB$ blood-type patients in the exchange pool. This implies that no triple with an $AB$ blood-type donor can be part of a 2-way exchange, since $AB$ blood-type donors can only donate to $AB$ blood-type patients.

We next argue that no triple with an $O$ blood-type patient can be part of a 2-way exchange. To see this, suppose that $X - Y - Z$ and $O - Y' - Z'$ take part in a 2-way exchange. If $X$ exchanges her $Y$ donor, then $Y$ can donate to $O$ so $Y = O$. If $X$ does not exchange her $Y$ donor, then $Y$ can donate to $X$. In either case, $Y \geq X$. Similarly $Z \geq X$, implying that $X - Y - Z$ is a compatible triple, a contradiction.

From what is shown above, the only triples that can be part of a 2-way exchange are those where the patient’s blood type is in \{A, B\} and the donors’ blood types are in \{O, A, B\}. If we further exclude the compatible combinations and repetitions due to reordering the donors, we are left with the six triple types stated in the Lemma. It is easy to verify that triples of these types can indeed participate in 2-way exchanges (see Figure 4).

Proof of Theorem 1: Let $N$ denote the maximum number of 2-way exchanges. Since each such exchange results in two transplants, the maximum number of transplants through 2-way exchanges is $2N$. We will prove the Theorem in two parts.

Proof of “$N \leq \min\{N_1, N_2, N_3, N_4\}$”: Since each 2-way exchange involves an $A$ blood-type patient, we have that $N \leq N_1$. Since $A - A - B$ types can only be part of a 2-way exchange with $B - B - A$ or $B - O - A$ types, the number of 2-way exchanges that involve an $A - A - B$ type is bounded above by $n(B - B - A) + n(B - O - A)$. Therefore, the number of 2-way exchanges involving an $A$ blood-type patient is less than or equal to this upper bound plus the number of $A - O - B$ and $A - B - B$ types, i.e., $N \leq N_2$. The inequalities $N \leq N_3$ and $N \leq N_4$ follow from symmetric arguments, switching the roles of $A$ and $B$ blood types.

Proof of “$N \geq \min\{N_1, N_2, N_3, N_4\}$”: We will next show that the matching algorithm achieves $\min\{N_1, N_2, N_3, N_4\}$ exchanges. This implies $N \geq \min\{N_1, N_2, N_3, N_4\}$. Since $N \leq \min\{N_1, N_2, N_3, N_4\}$, we conclude that $N = \min\{N_1, N_2, N_3, N_4\}$, and hence, the matching algorithm is optimal.

Case 1.\text{“}N_1 = \min\{N_1, N_2, N_3, N_4\}\text{”}: The inequalities $N_1 \leq N_2$, $N_1 \leq N_3$, and $N_1 \leq N_4$ imply that:

\[
\begin{align*}
n(A - A - B) &\leq n(B - B - A) + n(B - O - A) \\
n(A - B - B) &\leq n(B - A - A) + n(B - O - A) \\
n(A - A - B) + n(A - B - B) &\leq n(B - B - A) + n(B - A - A) + n(B - O - A)
\end{align*}
\]
Therefore, after the maximum number of $A - A - B$ and $B - B - A$ types and the maximum number of $A - B - B$ and $B - A - A$ types are matched in the first step, there are enough $B - O - A$ types to accommodate any remaining $A - A - B$ and $A - B - B$ types in the second step.

Since $N_1 \leq N_4$, there are at least $n(A - O - B)$ triples with $B$ blood-type patients who are not matched to $A - A - B$ and $A - B - B$ types in the first two steps. Therefore, all $A - O - B$ triples are matched to triples with $B$ blood-type patients in the second and third steps. The resulting matching involves $N_1$ exchanges, since all $A$ blood-type patients take part in a 2-way exchange.

**Case 2.** "$N_2 = \min\{N_1, N_2, N_3, N_4\}": Since $N_2 \leq N_1$, we have $n(A - A - B) \geq n(B - B - A) + n(B - O - A)$. Therefore, all $B - B - A$ types are matched to $A - A - B$ types in the first step. Similarly, $N_2 \leq N_4$ implies that $n(A - O - B) + n(A - B - B) \leq n(B - A - A)$. Therefore, all $A - B - B$ types are matched to $B - A - A$ types in the first step. In the second step, there are no remaining $B - B - A$ types, but there are enough $B - A - A$ types to accommodate all $A - O - B$ types. Similarly, in the second step, there are no remaining $A - B - B$ types, but there are enough $A - A - B$ types to accommodate all $B - O - A$ types. There are no more exchanges in the third step. The resulting matching involves $N_2$ 2-way exchanges.

The cases where $N_3$ and $N_4$ are the minimizers follow from symmetric arguments exchanging the roles of $A$ and $B$ blood types. ■

**Proof of Lemma 2:** As argued in the proof of Lemma 1, no $AB$ blood-type patient or donor can be part of a $K$-way exchange. Therefore, the only triples that can be part of a $K$-way exchange are those where its patient’s and its donors’ blood types are in $\{O, A, B\}$. After excluding the compatible combinations, we are left with the triple types listed above.

Take any $K$-way exchange. Since every triple type listed above has at least an $A$ or a $B$ blood-type donor, the $K$-way exchange involves an $A$ or a $B$ blood-type patient. If it involves an $A$ blood-type patient, then that patient brings in a $B$ blood-type donor, so it must also involve a $B$ blood-type patient. If it involves a $B$ blood-type patient, then that patient brings in an $A$ blood-type donor, so it must also involve an $A$ blood-type patient. It is trivial to see that all types in the hypothesis can feasibly participate in exchange in a suitable exchange pool.\(^{24}\) ■

**Proof of Lemma 4:** Suppose that there exists a 3-way exchange that matches an $O - X - X$ type triple for any $X \in \{A, B\}$. This triple’s $O$ patient necessarily receives grafts from two $O$ donors in this exchange. Then there exist two triples each with a single $O$ donor in the same exchange, as $X - O - O$ types are compatible and are not present in $\mathcal{E}$. By Lemma 2, there should be an $A$ and

\(^{24}\)We already demonstrated the possibility of exchanges regarding triples (of the types in the hypothesis of the lemma) with $A$ and $B$ blood type patients in Lemma 1. An $O - A - A$ triple can be matched in a four-way exchange with $A - O - B, A - O - B, B - A - A$ triples (symmetric argument holds for $O - B - B$). On the other hand, an $O - A - B$ triple can be matched in a 3-way exchange with $A - O - B$ and $B - O - A$ triples. An $O - O - A$ or an $O - O - B$ type can be used instead of $O - A - B$ in the previous example.
an \( B \) patient in any exchange. Thus, the other two triples that participate this 3-way exchange are necessarily of types \( A - O - B \) and \( B - O - A \), respectively (as types with \( AB \) patients or donors and \( A - O - O \), \( B - O - O \), \( A - O - A \), and \( B - O - B \) types do not participate in exchange by Lemma 2). However, \( O - X - X \), \( A - O - B \), and \( B - O - A \) types cannot form a feasible 3-way exchange among each other. This contradicts the existence of such a 3-way exchange. Hence, \( O - X - X \) types cannot participate in a 3-way exchange.

Before proving Theorem 2, we first state and prove two Lemmas that will be used in proving Theorem 2. Lemma 6 below states that, under the long-run assumption, one can restrict attention to the exchanges in Definition 3 to construct an optimal matching.

**Lemma 6** Suppose that the exchange pool \( \mathcal{E} \) satisfies the long-run assumption, and only 2- and 3-way exchanges are allowed. Then, there is an optimal matching that is in simplified form.

**Proof of Lemma 6:** We first show that if a matching \( \mu \) includes an exchange not represented in Figure 7, then there is a matching \( \mu' \) that induces at least as many transplants and includes one more exchange of the kinds included in Figure 7. To see this, take any exchange in \( \mu \) not represented as an edge in Figure 7. The exchange must be at most 3-way since larger exchanges are not allowed. Furthermore, by Lemma 2, the exchange includes two types \( A - Y - Z \) and \( B - Y' - Z' \) that are vertices of Figure 7. To create the matching \( \mu' \), we first undo this exchange in \( \mu \), then create a weakly larger exchange that involves unmatched types and is represented as an edge in Figure 7.

**Case 1.** “There is a bold edge between the types \( A - Y - Z \) and \( B - Y' - Z' \) in Figure 7”: Then we create the 3-way exchange that corresponds to that bold edge.

If there is no bold edge between \( A - Y - Z \) and \( B - Y' - Z' \) in Figure 7, then these types cannot be \( A - A - B \) and \( B - B - A \), because the only allowable exchange involving \( A - A - B \) and \( B - B - A \) is the 2-way exchange included in Figure 7. By an analogous argument, these types also cannot be \( A - B - B \) and \( B - A - A \). This leaves out two more cases:

**Case 2.** “\( A - Y - Z = A - A - B \) and \( B - Y' - Z = B - A - A \)”\: The only allowable exchange involving these two types not represented in Figure 7 is the 3-way exchange where the third participant is \( A - O - B \). In this case, we create the 3-way exchange that corresponds to the bold edge between the unmatched \( A - O - B \) and \( B - A - A \) types.

**Case 3.** “\( A - Y - Z = A - B - B \) and \( B - Y' - Z = B - B - A \)” \: We omit the argument for this case, since it is symmetric to Case 2.

By the finiteness of the problem, there is an optimal matching \( \mu \) that is not necessarily in simplified form. By what we have shown above, we can construct an optimal matching \( \mu' \) that is in simplified form from the matching \( \mu \) by iteratively replacing the exchanges that are excluded from Figure 7 with those that are included in it.
Proof of Theorem 2: Define the numbers $K_A$ and $K_B$ by:

\[ K_A := n(A - O - B) - n(B - B - A) - n(B - A - A) \]

Lemma 7 Suppose that the exchange pool $\mathcal{E}$ satisfies the long-run assumption and $n(A - A - B) + n(A - B - B) > n(B - O - A)$. If a matching $\mu$ is in simplified form and includes at least one 3-way exchange involving an $A - O - B$ and a $B - O - A$ type, then there is a matching $\mu'$ such that: (i) $\mu'$ is in simplified form, (ii) $\mu'$ induces at least as many transplants as $\mu$ and (iii) $\mu'$ includes one less 3-way exchange involving an $A - O - B$ and a $B - O - A$ type compared to $\mu$.

Proof of Lemma 7: To construct $\mu'$, we first undo exactly one 3-way exchange in $\mu$ that involves an $A - O - B$ and a $B - O - A$ type. In the following, we will call these $A - O - B$ and $B - O - A$ types, “the $A - O - B$ type” and “the $B - O - A$ type.” To finish constructing $\mu'$, we consider five cases:

Case 1. “There is an unmatched $A - A - B$ or $A - B - B$ type under $\mu$”: Then create a 3-way exchange involving that type and the $B - O - A$ type.

If we do not fall into Case 1, then all $A - A - B$ and $A - B - B$ types are matched under $\mu$; but since $n(A - A - B) + n(A - B - B) > n(B - O - A)$, they cannot all be part of a 3-way exchange with $B - O - A$ types. That leaves four more cases:

Case 2. “An $A - A - B$ and a $B - B - A$ type are part of a 2-way exchange under $\mu$”: Then undo that 2-way exchange and create two new 3-way exchanges, one involving the unmatched $A - A - B$ type and the $B - O - A$ type, and another involving the unmatched $B - B - A$ type and the $A - O - B$ type.

Case 3. “Two $A - A - B$ types and a $B - A - A$ type are part of a 3-way exchange under $\mu$”: Then undo that 3-way exchange and create two new 3-way exchanges, one involving one of the two unmatched $A - A - B$ types and the $B - O - A$ type, and another involving the unmatched $B - B - A$ type and the $A - O - B$ type.

Case 4. “An $A - B - B$ and a $B - A - A$ type are part of a 2-way exchange under $\mu$”: Then undo that 2-way exchange and create two new 3-way exchanges, one involving the unmatched $A - B - B$ type and the $B - O - A$ type, and another involving the unmatched $B - A - A$ type and the $A - O - B$ type.

Case 5. “An $A - B - B$ type and two $B - B - A$ types are part of a 3-way exchange under $\mu$”: Then undo that 3-way exchange and create two new 3-way exchanges, one involving the unmatched $A - B - B$ type and the $B - O - A$ type, and another involving one of the two unmatched $B - B - A$ types and the $A - O - B$ type.

In each of the five cases considered above, the newly constructed matching $\mu'$ satisfies (i)–(iii) in Lemma 7.

Proof of Theorem 2: Define the numbers $K_A$ and $K_B$ by:

\[ K_A := n(A - O - B) - n(B - B - A) - n(B - A - A) \]
\[ K_B := n(B - O - A) - n(A - A - B) - n(A - B - B) \]

We will consider two cases depending on the signs of \( K_A \) and \( K_B \).

**Case 1.** \( \max\{K_A, K_B\} \geq 0 \):

Suppose, without loss of generality, that \( K_A \leq K_B \). Then, \( K_B = \max\{K_A, K_B\} \geq 0 \). This implies, by the definition of \( K_B \), that \( n(B - O - A) \geq n(A - A - B) + n(A - B - B) \). Therefore, all \( A - A - B \) and \( A - B - B \) types participate in 3-way exchanges with \( B - O - A \) types in Step 2 of the algorithm.

The number of \( A - O - B \) types that are not matched in Step 2 is given by:

\[
\begin{align*}
n(A - O - B) - \min\{n(B - B - A) + n(B - A - A), n(A - O - B)\} \\
= \max\{n(A - O - B) - n(B - B - A) - n(B - A - A), 0\} \\
= \max\{K_A, 0\} \\
\leq K_B = n(B - O - A) - n(A - A - B) - n(A - B - B).
\end{align*}
\]

As a result, the number of \( A - O - B \) types that are not matched in Step 2 is less than or equal to the number of \( B - O - A \) types that are not matched in Step 2. Therefore, all \( A - O - B \) types participate in 3-way exchanges in Steps 2 and 3 of the algorithm.

We have shown that the algorithm creates at least \( 3 \times [n(A - A - B) + n(A - B - B) + n(A - O - B)] \) transplants. Since each exchange consists of at most three participants and must involve an \( A \) blood-type patient, this is also an upper bound on the number of transplants through 2- and 3-way exchanges. Therefore, the outcome of the algorithm must be optimal.

**Case 2.** \( \max\{K_A, K_B\} < 0 \):

By Lemma 6, there exists an optimal matching \( \mu_0 \) that is in simplified form. Since \( K_B < 0 \), we have \( n(A - A - B) + n(A - B - B) > n(B - O - A) \). Therefore, we can iteratively apply Lemma 7 to \( \mu_0 \) to obtain an optimal matching \( \mu_1 \) in simplified form that does not include a 3-way exchange involving an \( A - O - B \) and a \( B - O - A \) type.

Let \( \Delta_A \) denote the number of unmatched \( A - O - B \) types in \( \mu_1 \). Since \( K_A < 0 \), i.e., \( n(B - B - A) + n(B - A - A) > n(A - O - B) \), there are more than \( \Delta_A \) many participants with \( B - B - A \) or \( B - A - A \) types who do not take part in an exchange with \( A - O - B \) types in \( \mu_1 \). Choose an arbitrary \( \Delta_A \) many of these \( B - B - A \) or \( B - A - A \) participants, undo the exchanges they participate in under \( \mu_1 \), and create \( \Delta_A \) new 3-way exchanges involving these participants and the unmatched \( A - O - B \) types.

Similarly, let \( \Delta_B \) denote the number of unmatched \( B - O - A \) types in \( \mu_1 \). Since \( K_B < 0 \), i.e., \( n(A - A - B) + n(A - B - B) > n(B - O - A) \), there are more than \( \Delta_B \) many participants with \( A - A - B \) or \( A - B - B \) types who do not take part in an exchange with \( B - O - A \) types in \( \mu_1 \). Choose an arbitrary \( \Delta_B \) many of these \( A - A - B \) or \( A - B - B \) participants, undo the exchanges they participate in under \( \mu_1 \), and create \( \Delta_B \) new 3-way exchanges involving these participants and the unmatched \( B - O - A \) types.
The new matching $\mu_2$ obtained from $\mu_1$ in the above manner is in simplified form. Furthermore $\mu_2$ induces at least as many transplants as $\mu_1$; therefore, it is also optimal. Note also that under $\mu_2$, all $A-O-B$ types take part in a 3-way exchange with $B-B-A$ or $B-A-A$ types, and all $B-O-A$ types take part in a 3-way exchange with $A-A-B$ or $A-B-B$ types.

Let $\mu$ denote an outcome of the sequential matching algorithm described in the text. Since $K_A, K_B < 0$, the constraint (*) in Step 1 becomes equivalent to:

1. Leave at least a total $n(B-O-A)$ of $A-A-B$ and $A-B-B$ types unmatched.
2. Leave at least a total $n(A-O-B)$ of $B-B-A$ and $B-A-A$ types unmatched.

Therefore in Step 2 of the algorithm, all $A-O-B$ types take part in a 3-way exchange with $B-B-A$ or $B-A-A$ types, and all $B-O-A$ types take part in a 3-way exchange with $A-A-B$ or $A-B-B$ types. This implies that the total number of transplants from exchanges involving $A-O-B$ or $B-O-A$ types is the same ($= 3 \times [n(A-O-B) + n(B-O-A)]$) for both matchings $\mu_2$ and $\mu$.

The restriction of the matching $\mu_2$ to the 2- and 3-way exchanges represented as edges among $A-A-B$, $A-B-B$, $B-B-A$, and $B-A-A$ types in Figure 7 respects the constraint (*). Therefore, the total number of transplants in $\mu_2$ from exchanges not involving $A-O-B$ nor $B-O-A$ types cannot exceed the total number of transplants in Step 1 of the algorithm leading to $\mu$. As a result, the total number of transplants under $\mu$ is at least as large as the total number of transplants under $\mu_2$, implying that $\mu$ is also optimal.

Proof of Lemma 5: Suppose there are no exchange-size constraints and there exists an optimal matching $\mu$ that matches an $O-X-Y$ type triple $i$ where $X, Y \in \{A, B\}$ in $E$. By the long-run assumption, there exist an $O-O-X$ triple $j$ and an $O-O-Y$ triple $k$ that are unmatched in $\mu$. We construct a new matching $\nu$ using $\mu$ by removing triple $i$ and inserting triples $j$ and $k$ as follows (see Figure 10):

(i) the patient who originally receives from the $X$ donor of $i$ in $\mu$ now receives from the $X$ donor of $j$ in $\nu$,
(ii) the patient who originally receives from the $Y$ donor of $i$ in $\mu$ now receives from the $Y$ donor of $k$ in $\nu$,
(iii) the two $O$ donors who originally donate to the patient of $i$ in $\mu$ now donate to the patients of $j$ and $k$ in $\nu$, respectively,
(iv) the $O$ donors of $j$ and $k$ now donate to their own patients in $\nu$, and
(v) the other donations in $\mu$ remain intact in $\nu$.

The new matching $\nu$ is feasible and matches one more triple than $\mu$, which in turn contradicts the optimality of $\mu$. 

$\blacksquare$
Figure 10: The $O-X-Y$ triple in $\mu$ in the left is replaced with an $O-O-X$ triple and an $O-O-Y$ triple in $\nu$ in the right, which were originally unmatched in $\mu$. Since triples with $AB$ patients are absent in the exchange pool, the patients that $X$ and $Y$ donors donate in $\mu$ should also be of blood types $X$ and $Y$, respectively. Moreover, although it is depicted in the figure as if these donors donate to two different patients in $\mu$, when $X = Y$ it could also be the case that they donate to the same patient.

References


Appendix B  Dynamic Simulations

In the dynamic simulations, patients and their donors arrive over time and remain in the population until they are matched through exchange. We run statically optimal exchange algorithms once in each period.\textsuperscript{25} In each simulation, we generate $S = 500$ such populations and report the averages and sample standard errors of the simulation statistics.

B.1  Dynamic Simulations for Lung Exchange

In the dynamic lung-exchange simulations, we consider 200 triples arriving over 20 periods at a uniform rate of 10 triples per period. This time horizon roughly corresponds to more than 1 year of Japanese patient arrival, when exchanges are run once about every 3 weeks. We only consider the 2-way and 2&3-way exchange regimes.

Based on the 2&3-way exchange simulation results reported in Table 5, we can increase the number of living-donor transplants by 190%, thus nearly tripling them. This increase corresponds to 24% of all triples in the population. Even the logistically simpler 2-way exchange technology has a potential to increase the number of living-donor transplants by 125%.

\begin{center}
\begin{tabular}{llll}
\hline
\textbf{Population Size} & \textbf{Direct Donation} & \textbf{Exchange Tech.} \\
\hline
200 (in 20 periods) & 24.846 & 31.2 & 47.976 \\
\hline
& (4.5795) & (6.6568) & (8.7166) \\
\hline
\end{tabular}
\end{center}

Table 5: Dynamic lung-exchange simulations.

B.2  Dynamic Simulations for Dual-Graft Liver Exchange

For dual-graft liver exchange, we consider 500 triples arriving over 20 periods at a uniform rate of 25 triples per period. In each period, we follow the same 4-step transplantation scenario we used for the static simulations. The unmatched triples remain in the patient population waiting for the next period.\textsuperscript{26}

\textsuperscript{25}Moreover, among the optimal matchings, we choose a random one rather than using a priority rule to choose whom to match now and who to leave to future runs. The number of patients who can be matched dynamically can be further improved using dynamic optimization. For example, see Unver (2010) for such an approach for kidney exchanges.

\textsuperscript{26}Roughly, a dynamic simulation corresponds to 3.5 months in real time and the exchange is run once every 5 – 6 days. This is a very crude mapping that relies on our specific patient and paired donor generation assumptions. It
The dynamic simulation results are reported in Table 6. Under 2&3-way exchanges, the number of transplants via 2-donor exchange is only 15% short of those from 2-donor direct transplants, but 25% more than those from 1-donor exchanges. Hence dual-graft liver exchange is a viable modality under 2&3-way exchanges. With only 2-way exchanges, the number of transplants via 2-donor exchange is 39% less than those from 2-donor direct transplantation, but still 7% more than those from 1-donor exchange. In summary, dual-graft liver exchange increases the number of living-donor liver transplants by nearly 30% when 2&3-way exchanges are possible, and by more than 22% when only 2-way exchanges are possible.

### Dynamic Dual-Graft Liver-Exchange Simulations

<table>
<thead>
<tr>
<th>Population Size</th>
<th>1-Donor Direct</th>
<th>1-Donor Exchange</th>
<th>2-Donor Direct</th>
<th>2-Donor Exchange</th>
</tr>
</thead>
<tbody>
<tr>
<td>500 (in 20 periods)</td>
<td>119.83 (10.016)</td>
<td>58.284 (9.6765)</td>
<td>102.24 (10.05)</td>
<td>62.296 (10.016)</td>
</tr>
<tr>
<td>2&amp;3-way</td>
<td>68.034 (11.494)</td>
<td>100.47 (10.063)</td>
<td>85.632 (12.058)</td>
<td></td>
</tr>
</tbody>
</table>

**Table 6**: Dynamic dual-graft liver-exchange simulations.

### For Online Publication

#### Appendix C  Simultaneous Liver-Kidney Transplantation

For end-stage liver disease patients who also suffer from kidney failure, simultaneous liver-kidney transplantation (SLK) is a common procedure. In 2015, 626 patients received SLK transplants from deceased donors in the US. Transplanting a deceased-donor kidney to a (primarily liver) patient who lacks the highest priority on the kidney waitlist is a controversial topic, and this practice is actively debated in the US (Nadim et al., 2012). SLK transplants from living donors are not common in the US due to the low rate of living-donor liver donation. In contrast, living donation for both livers and kidneys is the norm in most Asian countries, such as South Korea, and countries with predominantly Muslim populations, such as Turkey. SLK transplantation from living donors is reported in the literature for these two countries, as well as for Azerbaijan and India (see Lee et al., 2011; Astarcioglu et al., 2003; Ahmadov et al., 2014; Aneja and Upwar, 2011, respectively).

For SLK exchange the analytical model we presented in Section 4 will be an approximation, since, in addition to the blood-type compatibility requirement of solid organs, kidney transplantation requires tissue-type compatibility and (single-graft) liver transplantation requires size compatibility. is obtained as follows: Under the current patient and donor generation scenario, a bit more than half of the patients will have at least one single-graft left- or right-lobe compatible donor or dual-graft compatible two donors (with more than 30% remnant donor liver). There are around 850 direct transplants a year in Korea from live donors, meaning that 1700 patients and their paired donors arrive a year. Then 500 patients arrive in roughly 3.5 months.
C.1 Simulations for Simultaneous Liver-Kidney Exchange

As our last application, we consider simulations for simultaneous liver-kidney (SLK) exchange. We use the underlying parameters reported in Table 3 in the main text and Table 7 here based on (mostly) Korean characteristics. In addition to an isolated SLK exchange, we also consider a possible integration of the SLK exchange with kidney-alone (KA) and liver-alone (LA) exchanges.

Statistics for SLK-Exchange Simulations
from the South Korean Kidney Patients/Donors

<table>
<thead>
<tr>
<th>Live Kidney Donation Recipients in 2010-2014</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Female</td>
<td>2555 (53.38%)</td>
</tr>
<tr>
<td>Male</td>
<td>2231 (46.62%)</td>
</tr>
<tr>
<td>Total</td>
<td>4786 (100.0%)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Live Kidney Donors in 2010-2014</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female</td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Patient PRA Distribution</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Range: 0-10%</td>
<td>70.19%</td>
</tr>
<tr>
<td>Range: 11-80%</td>
<td>20.00%</td>
</tr>
<tr>
<td>Range: 81-100%</td>
<td>9.81%</td>
</tr>
</tbody>
</table>

**Table 7:** Summary statistics for simultaneous-liver-kidney-exchange simulations, the kidney population. The patient PRA distribution is obtained from American UNOS data as we could not find detailed Korean PRA distributions. The transplant data were obtained from the Korean Network for Organ Sharing (KONOS) 2014 Annual Report, retrieved from https://www.konos.go.kr/konosis/index.jsp on 04/10/2016.

Following the South Korean statistics reported in these tables, we assume that the number of liver patients (LA and SLK) is \( \frac{9}{11} \)th of the number of kidney patients. We failed to find data on the percentage of South Korean liver patients who are in need of a SLK transplantation.\(^{27}\) Based on data from US, we consider two treatments where 7.5% and 15% of all liver patients are SLK candidates, respectively. We interpret these numbers as lower and upper bounds for SLK diagnosis prevalence.\(^{28}\)

We generate the patients and their attached donors as follows: We assume that each KA patient is paired with a single kidney donor, each LA patient is paired with a single liver donor, and each SLK patient is paired with one liver and one kidney donor. A kidney donor is deemed compatible with a kidney patient (KA or SLK) if he is blood-type and tissue-type compatible with the patient.\(^{29}\)

\(^{27}\) The gender of an SLK patient is determined using a Bernoulli distribution with a female probability as a weighted average of liver and kidney patients’ probabilities reported in Tables 3 and 7 with the ratio of weights 9 to 11.

\(^{28}\) In the US, according to the SLK transplant numbers given in Formica et al. (2016), 7.5% of all liver transplants involved SLK transplants between 2011-2015. On the other hand, Eason et al. (2008) report that only 73% of all SLK candidates received SLK transplants in 2006 and 2007 in the US. Moreover, Slack, Yeoman, and Wendon (2010) report that 47% of liver transplant patients develop either acute kidney injury (20%) or chronic kidney disease (27%), and patients from both of these categories could be suitable for SLK transplants.

\(^{29}\) For checking tissue-type compatibility, we generate a statistic known as panel reactive antibody (PRA) for
Following the methodology in Subsection 3.2, a liver donor is deemed compatible with a liver patient (LA or SLK) if he is blood-type compatible and his liver’s left lobe volume is at least 40% of the patient’s liver volume. An SLK patient participates in exchange if any one of her two donors are incompatible, and a KA or LA patient participates in exchange if her only donor is incompatible.

We consider two scenarios, referred to as “isolated” and “integrated” respectively, for our SLK simulations. For both scenarios, a kidney donor can be exchanged only with another kidney donor, and a liver donor can be exchanged only with another liver donor.

In the “isolated” scenario, we simulate the three exchange programs separately for each patient group, LA, KA, and SLK, using the 2-way exchange technology. Note that a 2-way exchange for the SLK group involves 4 donors, while a 2-way exchange for other groups involves 2 donors.

In the “integrated” scenario, we simulate a single exchange program to assess the potential welfare gains from a unification of individual exchange programs. For our simulations, we use the smallest meaningful exchange sizes that would fully integrate KA and LA with SLK. As such, we allow for any feasible 2-way exchange in our integrated scenario along with 3-way exchanges between one LA, one KA, and one SLK patient.

We consider two scenarios, referred to as “isolated” and “integrated” respectively, for our SLK simulations. For both scenarios, a kidney donor can be exchanged only with another kidney donor, and a liver donor can be exchanged only with another liver donor.

In the “isolated” scenario, we simulate the three exchange programs separately for each patient group, LA, KA, and SLK, using the 2-way exchange technology. Note that a 2-way exchange for the SLK group involves 4 donors, while a 2-way exchange for other groups involves 2 donors.

In the “integrated” scenario, we simulate a single exchange program to assess the potential welfare gains from a unification of individual exchange programs. For our simulations, we use the smallest meaningful exchange sizes that would fully integrate KA and LA with SLK. As such, we allow for any feasible 2-way exchange in our integrated scenario along with 3-way exchanges between one LA, one KA, and one SLK patient.

<table>
<thead>
<tr>
<th>SLK Patient Fraction in Liver Pool</th>
<th>Population Sizes</th>
<th>Direct Donation</th>
<th>Isolated Exchange Regime</th>
<th>Integrated Exchange Regime</th>
</tr>
</thead>
<tbody>
<tr>
<td>KA</td>
<td>SLK</td>
<td>LA</td>
<td>KA</td>
<td>SLK</td>
</tr>
<tr>
<td>7.5%</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>133</td>
<td>9</td>
<td>108</td>
<td>61.114</td>
<td>0.58</td>
</tr>
<tr>
<td>n = 250</td>
<td></td>
<td></td>
<td>5.944</td>
<td>0.70753</td>
</tr>
<tr>
<td>267</td>
<td>18</td>
<td>215</td>
<td>121.3</td>
<td>1.29</td>
</tr>
<tr>
<td>n = 500</td>
<td></td>
<td></td>
<td>8.3792</td>
<td>1.1119</td>
</tr>
<tr>
<td>535</td>
<td>35</td>
<td>430</td>
<td>244.09</td>
<td>2.426</td>
</tr>
<tr>
<td>15%</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>129</td>
<td>18</td>
<td>103</td>
<td>59.288</td>
<td>1.168</td>
</tr>
<tr>
<td>n = 250</td>
<td></td>
<td></td>
<td>5.90753</td>
<td>1.0421</td>
</tr>
<tr>
<td>259</td>
<td>36</td>
<td>205</td>
<td>117.64</td>
<td>2.566</td>
</tr>
<tr>
<td>n = 500</td>
<td></td>
<td></td>
<td>8.3432</td>
<td>1.5933</td>
</tr>
<tr>
<td>518</td>
<td>72</td>
<td>410</td>
<td>236.23</td>
<td>5.076</td>
</tr>
</tbody>
</table>

Table 8: Simultaneous liver-kidney exchange simulations for \( n = 250, 500, 1000 \) patients.

We consider population sizes of \( n = 250, 500, \) and \( 1000 \) for our simulations, reported in Table 8. For a population of \( n = 1000 \) and assuming 15% of liver patients are in need of SLK transplantation, the integrated exchange increases the number of SLK transplants over those from direct donation by 290%, almost quadrupling the number of SLK transplants. More than 20% of all SLK patients receive liver and kidney transplants through exchange in this case. For the same parameters, this exchange increases the number of SLK transplants over those from direct donation by 290%, almost quadrupling the number of SLK transplants. More than 20% of all SLK patients receive liver and kidney transplants through exchange in this case.

Each patient. PRA determines with what percentage of the general population the patient would have tissue-type incompatibility. The PRA distribution used in our simulations is reported in Table 7. Therefore, given the PRA value of a patient, we randomly determine whether a donor is tissue-type compatible with the patient.

We are not the first ones to propose a combined liver-and-kidney exchange. Dickerson and Sandholm (2014) show that higher efficiency can be obtained by combining kidney exchange and liver exchange if patients are allowed to exchange a kidney donor for a liver donor. Such an exchange, however, is quite unlikely given the very different risks associated with living-donor kidney donation and living-donor liver donation.
percentage reduces to 5.7% under the isolated scenario. Equivalently, the number of SLK transplants from an isolated SLK exchange is equal to 81% of the SLK transplants from direct transplantation. As such, integration of SLK with KA and LA increases transplants from exchange by about 260% for the SLK population.\(^\text{31}\)

When 7.5% of all liver patients are in need of SLK transplantation, integration becomes even more essential for the SLK patients. For a population of \(n = 1000\), exchange increases the number of SLK transplants by 55% under the isolated scenario. In contrast, exchange increases the number of SLK transplants by more than 300% under the integrated scenario. Hence, integration increases the number of transplants from exchange by almost 450%, matching 21% of all SLK patients.

C.2 Dynamic Simulations for Simultaneous Liver-Kidney Exchange

For simultaneous liver-kidney exchange dynamic simulations, we consider a population of \(n = 2000\) patients arriving over 20 periods under the identical regimes of our static simulations.\(^\text{32}\) Table 9 reports the results of these simulations.

<table>
<thead>
<tr>
<th>SLK Patient Population in Liver Pool</th>
<th>Direct Donation Isolated Sizes</th>
<th>Exchange Regime (n = 2000) Population</th>
<th>Direct Donation Integrated Sizes</th>
</tr>
</thead>
<tbody>
<tr>
<td>7.5% 1070 70 860 (in 20 periods)</td>
<td>KA SLK LA</td>
<td>KA SLK LA</td>
<td>KA SLK LA</td>
</tr>
<tr>
<td></td>
<td>487.84 4.946 134.93</td>
<td>306.62 5.588 110.35</td>
<td>312.92 18.246 113.17</td>
</tr>
<tr>
<td>15% 1036 144 820 (in 20 periods)</td>
<td>KA SLK LA</td>
<td>KA SLK LA</td>
<td>KA SLK LA</td>
</tr>
<tr>
<td></td>
<td>472.38 10.072 128.75</td>
<td>284.24 10.688 105.29</td>
<td>291.01 28.478 106.85</td>
</tr>
</tbody>
</table>

Table 9: Dynamic simultaneous liver-kidney exchange simulations for \(n = 2000\) patients.

When 15% of all liver patients are in need of SLK transplants, most outcomes essentially double with respect to the \(n = 1000\) static simulations: For LA and SLK, the changes are slightly more than 100%, while for KA the changes are slightly less than 100%. The increases for SLK patients are more substantial when 7.5% of all liver patients are in need of SLK transplants: An integrated exchange can facilitate transplants for 25% of all SLK patients, an overall increase of 360% with respect to SLK transplants from direct donation.

\(^{31}\)The contribution of integration is modest for other groups with an increase of 4% for KA transplants from exchange, and an increase of 4.5% for LA transplants from exchange.

\(^{32}\)This arrival rate roughly corresponds to 6 months of liver and kidney patients in South Korea with an exchange is carried out every 9 days.
Appendix D  The Subalgorithm for Algorithm 2

In this section, we present a subalgorithm that solves the constrained optimization problem in Step 1 of the matching algorithm for 2-\& 3-way exchanges. We define:

\[ \kappa_A := \min\{n(A - A - B) + n(A - B - B), n(B - O - A)\} \]
\[ \kappa_B := \min\{n(B - B - A) + n(B - A - A), n(A - O - B)\} \]

We can equivalently restate Step 1 by strengthening constraint (\(\ast\)) to be satisfied with equality:

Carry out the 2- and 3-way exchanges in Figure 7 among \(A - A - B\), \(A - B - B\), \(B - B - A\), and \(B - A - A\) types to maximize the number of transplants subject to the following constraints (\(\ast\ast\)):

1. Leave exactly a total of \(\kappa_A\) of \(A - A - B\) and \(A - B - B\) types unmatched.
2. Leave exactly a total of \(\kappa_B\) of \(B - B - A\) and \(B - A - A\) types unmatched.

\[ \text{Figure 11: The Exchanges in Step 1 of the 2-\& 3-way Matching Algorithm} \]

Figure 11 summarizes the 2-and 3-way exchanges that may be carried out in Step 1 above. In the following discussion, we restrict attention to the types and exchanges represented in Figure 11.

To satisfy the first part of constraint (\(\ast\ast\)), we can set aside any combination \(l_A\) of \(A - A - B\) types and \(m_A\) of \(A - B - B\) types, where \(l_A\) and \(m_A\) are integers satisfying

\[ 0 \leq l_A \leq n(A - A - B), \quad 0 \leq m_A \leq n(A - B - B), \quad \text{and} \quad l_A + m_A = \kappa_A. \]  

For any \(l_A\) and \(m_A\) satisfying Equation (1), the remaining number \(\gamma_A\) of \(B\) donors of \(A\) patients is:

\[ \gamma_A = n(A - A - B) - l_A + 2[n(A - B - B) - m_A] \tag{2} \]

Let \(\underline{l}_A\) and \(\overline{l}_A\) [\(\underline{m}_A\) and \(\overline{m}_A\)] be the smallest and largest values of \(l_A\) [\(m_A\)] among \((l_A, m_A)\) pairs that satisfy Equation (1). Then, the possible number of remaining \(B\) donors of \(A\) patients after
satisfying the first part of condition (**) is an integer interval \([\underline{\gamma}_A, \overline{\gamma}_A]\) where

\[
\gamma_A = n(A - A - B) - l_A + 2[n(A - B - B) - \bar{m}_A], \quad \text{and}
\]

\[
\overline{\gamma}_A = n(A - A - B) - \bar{l}_A + 2[n(A - B - B) - m_A].
\]

We can analogously define the integers \(l_B, \bar{l}_B, m_B, \bar{m}_B, \underline{\gamma}_B, \overline{\gamma}_B\) such that the possible number of remaining \(A\) donors of \(B\) patients that respect the second part of constraint (**) is an integer interval \([\underline{\gamma}_B, \overline{\gamma}_B]\).

In the first step of the subalgorithm, we determine which combination of types to set aside to satisfy constraint (**). We will consider three cases depending on the relative positions of the intervals \([\underline{\gamma}_A, \overline{\gamma}_A]\) and \([\underline{\gamma}_B, \overline{\gamma}_B]\).

**Subalgorithm 1 (Subalgorithm of the Sequential Matching Algorithm for 2- and 3-way Exchanges)**

**Step 1:**

We first determine \(\gamma_A\) and \(\gamma_B\):

**Case 1.** \(\{[\underline{\gamma}_A, \overline{\gamma}_A] \cap [\underline{\gamma}_B, \overline{\gamma}_B]\} \neq \emptyset\): Choose any \(\gamma_A = \gamma_B \in [\underline{\gamma}_A, \overline{\gamma}_A] \cap [\underline{\gamma}_B, \overline{\gamma}_B]\).

**Case 2.** \(\overline{\gamma}_A < \overline{\gamma}_B\):  

**Case 2.1.** If \(n(A - A - B) - \bar{l}_A - [n(B - B - A) - l_B]\) is positive and odd, and \(\underline{\gamma}_A < \gamma_A\), then set \(\gamma_A = \overline{\gamma}_A - 1\) and \(\gamma_B = \overline{\gamma}_B\).

**Case 2.2.** Otherwise, set \(\gamma_A = \overline{\gamma}_A\) and \(\gamma_B = \overline{\gamma}_B\).

**Case 3.** \(\overline{\gamma}_B < \overline{\gamma}_A\): Symmetric to Case 2, interchanging the roles of \(A\) and \(B\).

Then, we set aside \(l_A\) many \(A - A - B\)'s and \(m_A\) many \(A - B - B\)'s, where the integers \(l_A\) and \(m_A\) are uniquely determined by Equations (1) and (2) to ensure that the remaining number of \(B\) donors of \(A\) patients is \(\gamma_A\). The integers \(l_B\) and \(m_B\) are determined analogously.

**Step 2:**

In two special cases explained below, the second step of the subalgorithm sets aside one extra triple on top of those already set aside in Step 1.

**Case 1.** If \(\overline{\gamma}_A < \overline{\gamma}_B\), \(n(A - A - B) - \bar{l}_A - [n(B - B - A) - l_B]\) is positive and odd, \(\underline{\gamma}_A = \overline{\gamma}_A\), and \(n(B - B - A) - l_B > 0\), then set an extra \(B - B - A\) triple aside.

**Case 2.** If \(\overline{\gamma}_B < \overline{\gamma}_A\), \(n(B - B - A) - \bar{l}_B - [n(A - A - B) - l_A]\) is positive and odd, \(\underline{\gamma}_B = \overline{\gamma}_B\), and \(n(A - A - B) - \bar{l}_A > 0\), then set an extra \(A - A - B\) triple aside.

**Step 3:**
After having set the triples determined in Steps 1 and 2 of the subalgorithm aside, we sequentially maximize three subsets of exchanges among the remaining triples in Figure 11.

**Step 3.1:** Carry out the maximum number of 2-way exchanges between the $A-A-B$ and $B-B-A$ types.

**Step 3.2:** Carry out the maximum number of 3-way exchanges consisting of two $A-A-B$ and one $B-A-A$ triples, and those consisting of two $B-B-A$ and one $A-B-B$ triple, among the remaining types.

**Step 3.3:** Carry out the maximum number of 2-way exchanges between the remaining $A-B-B$ and $B-A-A$ types.

Figure 12 graphically illustrates the 2- and 3-way exchanges that are carried out at Steps 3.1–3.3 of the subalgorithm.

![Figure 12: Steps 3.1–3.3 of the Subalgorithm](image)

**Proposition 1** The subalgorithm described above solves the constrained optimization problem in Step 1 of the matching algorithm for 2- and 3-way exchanges.

**Proof of Proposition 1:** Constraint (*) is satisfied by construction, since in Step 1 of the subalgorithm $\gamma_i$ is chosen from $[\gamma_i^L, \bar{\gamma}_i]$ for $i = A, B$. Below, we show optimality by considering different cases.

**Case 1.** $[\bar{\gamma}_A, \hat{\gamma}_A] \cap [\bar{\gamma}_B, \hat{\gamma}_B] \neq \emptyset$: In this case, Step 1 of the subalgorithm sets $\gamma_A = \gamma_B$, i.e.:

$$n(A - A - B) - l_A + 2[n(A - B - B) - m_A] = n(B - B - A) - l_B + 2[n(B - A - A) - m_B]$$

and no extra triple is set aside in Step 2. Note that the above equality implies that at the end of Step 3.1 of the subalgorithm, the numbers of remaining $A - A - B$ and $B - B - A$ triples are even (at least one being zero). So again, by the above equality, all triples that are not set aside in Step 1 take part in 2- and 3-way exchanges by the end of Step 3 of the subalgorithm. This implies optimality.
Case 2. “$\bar{\gamma}_A < \underline{\gamma}_A$, i.e.,

$$n(A - A - B) - \bar{l}_A + 2[n(A - B - B) - m_A] < n(B - B - A) - \bar{l}_B + 2[n(B - A - A) - \bar{m}_B]$$ \hspace{1cm} (3)$$

We next establish an upper bound on the number of triples with $B$ patients that can participate in 2- and 3-way exchanges. Suppose that $p_B$ many $B - B - A$ triples and $r_B$ many $B - A - A$ triples can take part in 2- and 3-way exchanges while respecting condition (*). Since matching each $B - B - A$ triple requires one $B$ donor of an $A$ patient; matching each $B - A - A$ triple requires two $B$ donors of $A$ patients; and the maximum number of $B$ donors of $A$ patients is $\bar{\gamma}_A$; we have the constraint:

$$p_B + 2r_B \leq \bar{\gamma}_A$$

Note also that $p_B \leq \bar{p}_B := n(B - B - A) - \bar{l}_B$. Therefore, we cannot match any more triples with $B$ patients than the bound:

$$\bar{p}_B + \frac{1}{2}(\bar{\gamma}_A - \bar{p}_B) = \max_{p_B,r_B \in \mathbb{R}} \text{ s.t. } p_B + 2r_B \leq \bar{\gamma}_A$$

$$p_B \leq \bar{p}_B$$ \hspace{1cm} (4)$$

Case 2.1. “$n(A - A - B) - \bar{l}_A - [n(B - B - A) - \bar{l}_B] \text{ is positive and odd, and } \underline{\gamma}_A < \bar{\gamma}_A$”:

Note that $\gamma_A = \bar{\gamma}_A - 1$ and $\gamma_B = \bar{\gamma}_B$ imply that $l_A = \bar{l}_A - 1$, $m_A = m_A + 1$, $l_B = \bar{l}_B$, and $m_B = \bar{m}_B$. So $n(A - A - B) - l_A - [n(B - B - A) - \bar{l}_B]$ is positive and even. Furthermore, no extra triple is set aside in Step 2. Therefore, an even number of $A - A - B$ types remain unmatched at the end of Step 3.1. Also, by Equation (3),

$$n(A - A - B) - l_A + 2[n(A - B - B) - m_A] < n(B - B - A) - l_B + 2[n(B - A - A) - m_B].$$ \hspace{1cm} (5)$$

So all the $A - B - B$ types available at the end of Step 3.1 take part in 3-way exchanges with $B - A - A$ types in Step 3.2, and there are enough remaining $B - A - A$ types to accommodate all $A - B - B$ types in Step 3.3. Therefore, all triples with $A$ donors that are not set aside in Step 1 take part in 2- and 3-way exchanges in Step 3 of the subalgorithm.

We next show that it is impossible to match more triples with $B$ patients while respecting constraint (*), which will prove optimality. Since in Case 2.1, $\bar{\gamma}_A - \bar{p}_B$ is odd, and $\gamma_A = \bar{\gamma}_A - 1$, rounding down the upper bound in Equation (4) to the nearest integer gives:

$$\bar{p}_B + \frac{1}{2}(\bar{\gamma}_A - \bar{p}_B).$$

Note that this is the number of triples with $B$ patients who take part in 2- and 3-way exchanges in Step 3 of the subalgorithm. (In Step 3.1, $\bar{p}_B \equiv n(B - B - A) - \bar{l}_B$ many $B - B - A$ triples take part in 2-way exchanges; and in Steps 3.2 and 3.3, $\frac{1}{2}(\bar{\gamma}_A - \bar{p}_B)$ many $B - A - A$ triples take part in 2- and 3-way exchanges.)
Case 2.2. We further break Case 2.2 into four subcases:

Case 2.2.1. \( n(A - A - B) - \tilde{l}_A - [n(B - B - A) - \bar{l}_B] \) is positive and odd, \( \gamma_A = \bar{\gamma}_A \), and \( n(B - B - A) - \bar{l}_B > 0 \):

Note that \( \gamma_A = \bar{\gamma}_A \) and \( \gamma_B = \bar{\gamma}_B \) imply that \( l_A = \tilde{l}_A, m_A = \bar{m}_A, l_B = \bar{l}_B, \) and \( m_B = \bar{m}_B \). So \( n(A - A - B) - l_A - [n(B - B - A) - l_B] \) is positive and odd, and Equation (5) holds. Since one more \( B - B - A \) triple is set aside in Step 2, an even number of \( A - A - B \) types remain unmatched at the end of Step 3.1. By Equation (5), all triples with \( A \) donors that are not set aside in Step 1 take part in 2- and 3-way exchanges in Step 3 of the subalgorithm.

We next show that it is impossible to match more triples with \( B \) patients while respecting constraint (*), which will prove optimality. Since in this case, \( \bar{\gamma}_A - \bar{p}_B \) is odd and \( \gamma_A = \bar{\gamma}_A \), rounding down the upper bound in Equation (4) to the nearest integer gives:

\[
\bar{p}_B - 1 + \frac{1}{2}[\gamma_A - (\bar{p}_B - 1)].
\]

Note that this is the number of triples with \( B \) patients who take part in 2- and 3-way exchanges in Step 3 of the subalgorithm. (In Step 3.1, \( \bar{p}_B - 1 \equiv n(B - B - A) - \bar{l}_B - 1 \) many \( B - B - A \) triples take part in 2-way exchanges; and in Steps 3.2 and 3.3, \( \frac{1}{2}[\gamma_A - (\bar{p}_B - 1)] \) many \( B - A - A \) triples take part in 2- and 3-way exchanges.)

Case 2.2.2. \( n(A - A - B) - \tilde{l}_A - [n(B - B - A) - \bar{l}_B] \) is positive and odd, \( \gamma_A = \bar{\gamma}_A \), and \( n(B - B - A) - \bar{l}_B = 0 \):

Since \( n(B - B - A) \geq \tilde{l}_B \geq \bar{l}_B \) and \( n(B - B - A) - \bar{l}_B = 0 \), we have \( \tilde{l}_B = \bar{l}_B \), which implies that \( \gamma_B = \bar{\gamma}_B \). Since \( \gamma_A = \bar{\gamma}_A \) and \( \gamma_B = \bar{\gamma}_B \) in this case, the choices of \( \gamma_A \) and \( \gamma_B \) in Step 1 of the subalgorithm correspond to the unique way of satisfying constraint (**). That is, \( \gamma_A = \gamma_B = \bar{\gamma}_A \) and \( \gamma_B = \gamma_B = \bar{\gamma}_B \), \( l_A = \tilde{l}_A, m_A = \bar{m}_A, l_B = \bar{l}_B, \) and \( m_B = \bar{m}_B \). Also, Equation (5) holds.

So \( n(A - A - B) - \tilde{l}_A \) is positive and odd, and \( n(B - B - A) - \bar{l}_B = 0 \). Furthermore, no extra triple is set aside in Step 2. Therefore, there are no matches in Step 3.1 and all of the (odd number of) \( A - A - B \) triples are available in the beginning of Step 3.2. By Equation (5), all but one of these \( A - A - B \) triples take part in 3-way exchanges with \( B - A - A \) types in Step 3.2; and there are enough remaining \( B - A - A \) types to accommodate all \( A - B - B \) types in Step 3.3. Therefore, all triples with \( A \) donors, except one \( A - A - B \) triple, that are not set aside in Step 1 take part in 2- and 3-way exchanges in Step 3 of the subalgorithm.

To see that it is not possible to match any more triples with \( A \) patients, remember that in the current case the combination of triples that are set aside in Step 1 of the algorithm is determined uniquely; and note that since there are no remaining \( B - B - A \) triples, the \( A - A - B \) triples can only participate in 3-way exchanges with \( B - A - A \) triples. Each such 3-way exchange requires exactly two \( A - A - B \) triples; therefore, it is impossible to match all of the (odd number of) \( A - A - B \) triples.
We next show that it is impossible to match more triples with $B$ patients while respecting constraint (*), which will prove optimality. Since in Case 2.2.2, $\bar{\gamma}_A \leq \bar{p}_B$ is odd and $\gamma_A = \bar{\gamma}_A$, rounding down the upper bound in Equation (4) to the nearest integer gives:

$$\bar{p}_B + \frac{1}{2}[(\gamma_A - 1) - \bar{p}_B].$$

Note that this is the number of triples with $B$ patients who take part in 2- and 3-way exchanges in Step 3 of the subalgorithm. (In Step 3.1, $\bar{p}_B \equiv n(B - B - A) - l_B$ many $B - B - A$ triples take part in 2-way exchanges; and in Steps 3.2 and 3.3, $\frac{1}{2}[(\gamma_A - 1) - \bar{p}_B]$ many $B - A - A$ triples take part in 2- and 3-way exchanges.)

**Case 2.2.3.** "$n(A - A - B) - \bar{l}_A - [n(B - B - A) - l_B]$ is positive and even":

Note that $\gamma_A = \bar{\gamma}_A$ and $\gamma_B = \bar{\gamma}_B$ imply that $l_A = \bar{l}_A$, $m_A = \bar{m}_A$, $l_B = l_B$, and $m_B = \bar{m}_B$. So $n(A - A - B) - l_A - [n(B - B - A) - l_B]$ is positive and even and Equation (5) holds. Since no other triple is set aside in Step 2, an even number of $A - A - B$ types remain unmatched at the end of Step 3.1. By Equation (5), all triples with $A$ donors that are not set aside in Step 1 take part in 2- and 3-way exchanges in Step 3 of the subalgorithm.

We next show that it is impossible to match more triples with $B$ patients while respecting constraint (*), which will prove optimality. Since in this case, $\bar{\gamma}_A - \bar{p}_B$ is even and $\gamma_A = \bar{\gamma}_A$, the upper bound in Equation (4) is integer valued:

$$\bar{p}_B + \frac{1}{2}[\gamma_A - \bar{p}_B].$$

Note that this is the number of triples with $B$ patients who take part in 2- and 3-way exchanges in Step 3 of the subalgorithm. (In Step 3.1, $\bar{p}_B \equiv n(B - B - A) - l_B$ many $B - B - A$ triples take part in 2-way exchanges; and in Steps 3.2 and 3.3, $\frac{1}{2}(\gamma_A - \bar{p}_B)$ many $B - A - A$ triples take part in 2- and 3-way exchanges.)

**Case 2.2.4.** "$n(A - A - B) - \bar{l}_A - [n(B - B - A) - l_B] \leq 0$":

Note that $\gamma_A = \bar{\gamma}_A$ and $\gamma_B = \bar{\gamma}_B$ imply that $l_A = \bar{l}_A$, $m_A = \bar{m}_A$, $l_B = l_B$, and $m_B = \bar{m}_B$. Also Equation (5) holds. Since no other triple is set aside in Step 2 and $n(B - B - A) - l_B \geq n(A - A - B) - l_A$, all $A - A - B$ triples are matched in Step 3.1. By Equation (5), there are sufficient remaining $B - B - A$ and $B - A - A$ triples to ensure that all $A - B - B$ triples take part in 2- and 3-way exchanges in Steps 3.2 and 3.3. So all triples with $A$ donors that are not set aside in Step 1 take part in 2- and 3-way exchanges in Step 3 of the subalgorithm.

We next show that it is impossible to match more triples with $B$ patients while respecting constraint (*) by considering three cases, which will prove optimality.

Suppose first that $\bar{\gamma}_A - \bar{p}_B \leq 0$. Since matching each triple with a $B$ patient requires at least one $B$ donor of an $A$ patient and the maximum number of $B$ donors of $A$ patients is $\bar{\gamma}_A$, we cannot match more triples with a $B$ patient than $\bar{\gamma}_B$. Since in this case $n(B - B - A) - \bar{l}_B \equiv \bar{p}_B \geq \bar{\gamma}_A = \gamma_A$, 

43
the subalgorithm matches $\bar{\gamma}_A$ many $B - B - A$ triples in Steps 3.1 and 3.2, which achieves this upper bound.

Suppose next that $\bar{\gamma}_A - \bar{p}_B$ is positive and even. Then, the upper bound in Equation (4) is integer valued, and since $\gamma_A = \bar{\gamma}_A$, it can be written as:

$$\bar{p}_B + \frac{1}{2}(\gamma_A - \bar{p}_B).$$

Note that this is the number of triples with $B$ patients who take part in 2- and 3-way exchanges in Step 3 of the subalgorithm. (In Steps 3.1 and 3.2, $\bar{p}_B \equiv n(B - B - A) - l_B$ many $B - B - A$ triples take part in 2- and 3-way exchanges; and in Step 3.3, $\frac{1}{2}(\gamma_A - \bar{p}_B)$ many $B - A - A$ triples take part in 2-way exchanges.)

Suppose last that $\bar{\gamma}_A - \bar{p}_B$ is positive and odd. Then, since $\gamma_A = \bar{\gamma}_A$, rounding down the upper bound in Equation (4) to the nearest integer gives:

$$\bar{p}_B - 1 + \frac{1}{2}[\gamma_A - (\bar{p}_B - 1)].$$

Note that this is the number of triples with $B$ patients who take part in 2- and 3-way exchanges in Step 3 of the subalgorithm. (In Steps 3.1 and 3.2, $\bar{p}_B - 1 \equiv n(B - B - A) - l_B - 1$ many $B - B - A$ triples take part in 2- and 3-way exchanges; and in Step 3.3, $\frac{1}{2}[\gamma_A - (\bar{p}_B - 1)]$ many $B - A - A$ triples take part in 2-way exchanges.)

Case 3. “$\bar{\gamma}_B < \gamma_A$”: Symmetric to Case 2, interchanging the roles of $A$ and $B$.

For Online Publication

Appendix E  Proof of Theorem 3 and Other Results for Unrestricted Exchanges

Before delving into the analysis, we introduce some new terminology. For a given exchange pool $\mathcal{E}$, we refer to an exchange pool $\mathcal{K} \leq \mathcal{E}$ as a subpool of $\mathcal{E}$. We fix a dual-donor exchange pool $\mathcal{E}$ throughout the section. Given a subpool $\mathcal{K}$ let $D_X[\mathcal{K}]$ be the number of $X$ blood-type donors in $\mathcal{K}$ and $P_X[\mathcal{K}]$ as the number of $X$ blood-type patients in $\mathcal{K}$. We also use $n(X - Y - Z)[\mathcal{K}]$ to denote the number of $X - Y - Z$ triples in $\mathcal{K}$ (while we omit the arguments of these expressions if $\mathcal{K} = \mathcal{E}$). For a subpool $\mathcal{K}$, by a slight abuse of notation, let $|\mathcal{K}|$ be the total number of triples in $\mathcal{K}$. Given a matching $\mu$, we will sometimes denote the subpool of triples matched through it also as $\mu$, with a slight abuse of notation.
We denote with $\mathcal{E}$ the essential types:


Recall that by Lemma 2 each exchange should have at least two pairs of two different types in $\mathcal{E}$, one with an $A$ patient and one with a $B$ patient. Let $\mathcal{E}_E \leq \mathcal{E}$ be the subpool with only essential-type triples.

Also recall that by Lemma 5, the only types that can be a part of an optimal matching in this setting besides the essential types are $O - O - A$ and $O - O - B$, which are sufficiently many by the lung-run assumption. The following lemma will characterize the role of such triples and reduce the problem to focus only on essential types in constructing an optimal matching. This is a counterpart of Lemma 6 for unrestricted exchange sizes:

**Lemma 8** Suppose that $\mathcal{E}$ satisfies the long-run assumption and $\mu$ is an optimal matching in the absence of any exchange-size constraints in the essential type subpool $\mathcal{E}_E$. Suppose further that $\mu$ matches the maximum possible number of $A - O - B$ and $B - O - A$ triples that can be matched in any matching in $\mathcal{E}_E$.

1. Then $\mu$ can be modified to obtain a matching $\nu$ such that $n(A - O - B)[\mu] + n(B - O - A)[\mu]$-many $O - O - A$ and $O - O - B$ triples can be matched in addition to all triples matched by $\mu$.

2. Moreover, $\nu$ is an optimal matching of $\mathcal{E}$ in the absence of any exchange-size constraints.

**Proof of Lemma 8:** The first part of the Lemma is easy to prove: Take any $B - O - A$ or $A - O - B$ triple $i$ matched in $\mu$. Observe that as $\mu$ is a matching within essential types, the $O$ donor of $i$ is necessarily donating to an $A$ or a $B$ patient in $\mu$. If $i$’s $O$ donor is donating to an $A$ patient then take a triple $j$ of type $O - O - A$ and otherwise take a triple $j$ of type $O - O - B$. Such triples (which are unmatched in $\mu$) exist by the long-run assumption. Modify $\mu$ as follows: Let $i$’s and $j$’s $O$ donors donate to $j$’s patient and $j$’s non-$O$ donor donate to the patient $i$’s $O$ donor was previously donating to in $\mu$. Otherwise, do not change any other donations in $\mu$. We repeat the procedure for all $B - O - A$ and $A - O - B$ triples matched in $\mu$. Let $\nu$ be the matching obtained as a result of this procedure. It matches $n(A - O - B)[\mu] + n(B - O - A)[\mu]$-many $O - O - A$ and $O - O - B$ triples.

For the second part of the Lemma, we first prove a claim:

**Claim:** For any optimal matching $\nu'$ of $\mathcal{E}$, we can construct another matching $\mu'$ involving only the essential-type triples matched by $\nu'$.

**Proof of the Claim:** By Lemma 5, besides the essential-type triples, $O - O - A$ and $O - O - B$ types can participate in $\nu'$. Take a patient of a triple matched in $\nu'$ of type $O - O - X$ for any $X \in \{A, B\}$.
Without loss of generality assume that her $O$ donor directly donates to her in $\nu'$. Another triple’s $O$ donor $d_1$ donates to her in $\nu'$ as well. In return, her $X$ donor donates to a (different) patient $p_1$ in $\nu'$. We can simply take this $O - O - X$ triple out, and form a new matching by $d_1$ directly donating to $p_1$ and rest of the donations remain intact as in $\nu'$. We repeat this procedure for all triples of types $O - A - B$, $O - A - A$, $O - B - B$, $O - O - A$, and $O - O - B$ iteratively. The final matching, we refer to as $\mu'$, is feasible and consists of only essential-type triples of $\nu'$. □

Let $\mu$ and $\nu$ be defined as in the hypothesis of the Lemma. Suppose that $\nu'$ is an arbitrary optimal matching in $E$. We will show that $|\nu| = |\nu'|$.

By Lemma 5, the types of triples that can be part of a feasible exchange besides the essential types are $O - O - A$ and $O - O - B$ under the long-run assumption.

We form a matching $\mu'$ by removing the non-essential-type triples from $\nu'$ by the Claim. We have

$$|\mu| \geq |\mu'|$$ \hspace{1cm} (6)

by optimality of $\mu$ in $E$. We also have

$$n(A - O - B)[\mu] + n(B - O - A)[\mu] \geq n(A - O - B)[\mu'] + n(B - O - A)[\mu']$$ \hspace{1cm} (7)

by the fact that $\mu$ maximizes the number of $A - O - B$ and $B - O - A$ triples matched in $E$.

The $O - O - A$ and $O - O - B$ triples matched in $\nu'$ require at least $n(O - O - B)[\nu'] + n(O - O - A)[\nu']$-many other triples with $O$ donors in $\nu'$. Since these triples can only be of types $A - O - B$ and $B - O - A$, we have

$$n(O - O - A)[\nu] + n(O - O - B)[\nu] = n(A - O - B)[\mu] + n(B - O - A)[\mu]$$
$$\geq n(A - O - B)[\mu'] + n(B - O - A)[\mu']$$
$$\geq n(O - O - A)[\nu'] + n(O - O - B)[\nu']$$,

where the equality follows from the construction of $\nu$, the first inequality follows from Equation 7, and the last inequality follows from the feasibility of $\nu'$. This and Equation 6 imply $|\nu| = |\mu| + n(O - O - A)[\nu] + n(O - O - B)[\nu] \geq |\mu'| + n(O - O - A)[\nu'] + n(O - O - B)[\nu'] = |\nu'|$. Since $\nu'$ is optimal in $E$, $\nu$ is optimal in $E$ with $|\nu| = |\nu'|$, completing the proof. ■

If we can show that it is possible to construct a matching $\mu$, which simultaneously matches

1. the maximum number of $A - O - B$ and $B - O - A$ triples in any possible matching, and
2. the maximum number of essential-type triples,
then using Lemma 8, we can construct an optimal matching using \( \mu \), and it matches \( |\mu| + n(A - O - B)|\mu| + n(B - O - A)|\mu| \) triples. This will also give us the optimal number of triples that can be matched in the absence of exchange-size constraints.

Hence, our larger goal is to reach the above two goals simultaneously. Next, we define two non-negative numbers for triples in \( E \). These tell us the minimum (\( \underline{s}_A \)) and maximum (\( \bar{s}_A \)) numbers of donors compatible with \( B \) blood-type patients that can be supplied by patients with \( A \) blood-type patients:

\[
\begin{align*}
\underline{s}_A &:= n(A - O - B) + n(A - A - B) + 2n(A - B - B) \quad (8) \\
\bar{s}_A &:= 2n(A - O - B) + n(A - A - B) + 2n(A - B - B) \quad (9)
\end{align*}
\]

Here, \( \underline{s}_A \) assumes that all \( A - O - B \) triples are treated like \( A - A - B \) types: the \( O \) blood-type donor can be utilized internally, and hence, each \( A - O - B \) triple requires one donor from outside, and so does each \( A - A - B \) triple. On the other hand, each \( A - B - B \) triple needs two donors from outside.

In calculating \( \bar{s}_A \), we treat \( A - O - B \) triples like \( A - B - B \)’s. Therefore, each of them requires two donors from outside instead of one. Symmetrically, we define \( \underline{s}_B \) and \( \bar{s}_B \). Observe that

\[
\bar{s}_A - \underline{s}_A = n(A - O - B) \quad \text{and} \quad \bar{s}_B - \underline{s}_B = n(B - O - A).
\]

We define a subalgorithm using these numbers:


**Group:** Two cases are possible for \( \underline{s}_A, \bar{s}_A, \underline{s}_B, \bar{s}_B \), defined in Equations 8 and 9.

**Case 1.** \( \{[\underline{s}_A, \bar{s}_A] \cap [\underline{s}_B, \bar{s}_B] \neq \emptyset \} \):

Fix \( \alpha_A, \alpha_B \) such that \( 0 \leq \alpha_A \leq n(A - O - B), 0 \leq \alpha_B \leq n(B - O - A) \), and \( \bar{s}_A - \alpha_A = \bar{s}_B - \alpha_B \):

1. **Group** \( \alpha_A \)-many \( A - O - B \) triples with \( A - A - B \) types and the rest with \( A - B - B \) types.
2. **Group** \( \alpha_B \)-many \( B - O - A \) triples with \( B - B - A \) types and the rest with \( B - A - A \) types.

**Case 2.** \( \bar{s}_B < \underline{s}_A \):

1. **Group** all \( A - O - B \) triples (that is, \( (\bar{s}_A - \underline{s}_A) \)-many) with \( A - A - B \) types (i.e., \( \alpha_A = \bar{s}_A - \underline{s}_A \)).
2. **Group** all \( B - O - A \) triples (that is \( (\bar{s}_B - \underline{s}_B) \)-many) with \( B - A - A \) types (i.e., \( \alpha_B = \bar{s}_B - \underline{s}_B \)).
Case 3. $\bar{s}_A < \bar{s}_B$: Symmetric situation with Case 2 replacing $A$ blood type with $B$.

We refer to all $X-O-Z$ triples grouped with $X-Y-Z$ triples and all $X-Y-Z$ triples for all for $X,Y,Z \in \{A,B\}$ such that $X \neq Z$ as $X-Y^* - Z$ triples or group. Let $n(X-Y^* - Z)$ be the number of triples in $X-Y^* - Z$ group and $n(X-O-Z|X-Y^* - Z)$ be the number of $X-O-Z$ triples in the $X-Y^* - Z$ group. Define:

$$\Delta := n(A - A^* - B) - n(B - B^* - A).$$

(10)

**Match:** Starting with the triples with $O$ donors in each group in each step:

**Step 1:** Carry out the maximum number of 2-way exchanges between the $A - A^* - B$ and $B - B^* - A$ triples with the following exceptions:

(I) In Case 2 if $\Delta < 0$ and is odd:

(A) If $n(B - O - A) > 0$:

add one $B - O - A$ triple to $B - B^* - A$ group from $B - A^* - A$ group and continue with Step 1.

(B) If $n(B - O - A) = 0$ and an exchange can be conducted in Step 1: do not conduct the last $A - A^* - B & B - B^* - A$ 2-way exchange in Step 1 (and thus, exactly one $A - A^* - B$ triple and an even number of $B - B^* - A$ triples remain unmatched in Step 1), and if the remaining $A - A^* - B$ triple is of type $A - O - B$, then move it to the $A - B^* - B$ group.

(II) In Case 3 if $\Delta > 0$ and is odd:

(A) If $n(A - O - B) > 0$:

add one $A - O - B$ triple to $A - A^* - B$ group from $A - B^* - B$ group and continue with Step 1.

(B) If $n(A - O - B) = 0$ and an exchange can be conducted in Step 1:

do not conduct the last $A - A^* - B & B - B^* - A$ 2-way exchange in Step 1 (and thus, exactly one $B - B^* - A$ triple and an even number of $A - A^* - B$ triples remain unmatched in Step 1), and if the remaining $B - B^* - A$ triple is of type $B - O - A$, then move it to the $B - A^* - A$ group.

**Step 2:** Carry out the maximum number of 3-way exchanges consisting of two $A - A^* - B$ triples and one $B - A^* - A$ triple, and those consisting of two $B - B^* - A$ triples and one $A - B^* - B$ among the remaining ones.

**Step 3:** If there are any $A - O - B$ and $B - O - A$ triples left in $A - A^* - B$ and $B - B^* - A$ groups, respectively, then move them to $A - B^* - B$ and $B - A^* - A$ groups, respectively. Carry out the maximum number of 2-way exchanges between the remaining $A - B^* - B$ and $B - A^* - A$ triples.

Figure 13 summarizes how the Group and Match Subalgorithm works, along with its consequences (to be proven in Propositions 2 and 3 below). This subalgorithm is embedded in the
optimal matching algorithm as follows:

Algorithm 3 (Sequential Matching Algorithm in the Absence of Exchange-Size Constraints)

Step 1: Use Subalgorithm 2, Group and Match, to match triples of types $E$.

Step 2: In any exchange determined in this matching, for each $A-O-B$ or $B-O-A$ triple in the exchange, insert an $O-O-A$ or an $O-O-B$ triple using Lemma 8.

Before proving the optimality of Algorithm 3, we find an upper bound to the number of triples that can be matched in an exchange pool:

Lemma 9 (Upper Bound Lemma) Consider the subpool $E_E$. Then $\overline{m}$, defined below, is an upper
bound to the number of triples that can be matched in a matching consisting only of triples in $E_E$:

$$\overline{m} := \overline{m}_A + \overline{m}_B \text{ where}$$

$$\overline{m}_A := \min \left\{ P_A[E_E], \left\lfloor \frac{D_A[E_E] + D_O[E_E]}{2} \right\rfloor, \overline{s}_B \right\}$$

and

$$\overline{m}_B := \min \left\{ P_B[E_E], \left\lfloor \frac{D_B[E_E] + D_O[E_E]}{2} \right\rfloor, \overline{s}_A \right\}.$$

**Proof of Lemma 9:** The first term in $\overline{m}_A$, $P_A[E_E]$, is the number of $A$ blood-type patients and the second term, $\left\lfloor \frac{D_A[E_E] + D_O[E_E]}{2} \right\rfloor$, is the maximum number of $A$ blood-type patients who can receive two lobes from donors who are compatible with $A$ blood-type patients, i.e., $O$ and $A$ blood-type donors in $E_E$. Hence, each of them is an upper bound for the number of triples with $A$ blood-type patients in $E_E$ who can receive transplant. Next consider the third term: $\overline{s}_B = n(B - B - A) + 2n(B - O - A) + 2n(B - A - A)$ is the maximum number of $A$ or $O$ blood-type donors whom the $B$ blood-type patients can provide for the triples with $A$ blood-type patients in $E_E$. Each triple with an $A$ blood-type patient in $E_E$ requires at least one $A$ or $O$ blood-type donor coming from another triple to be matched feasibly, as it can provide at most one compatible donor for itself. Hence, $\overline{s}_B$ is also an upper bound to the number of $A$ blood-type patients who can be matched within $E_E$, establishing the formula for $\overline{m}_A$.

The argument is the same in $\overline{m}_B$ for $B$ blood-type patients. There are no triples with $AB$ or $O$ blood-type patients in $E_E$. This concludes the proof and establishes $\overline{m}$ as an upper bound to the essential types that can be matched.

We will prove that the upper bound found above is almost tight, and Group and Match subalgorithm matches always at least one fewer patient than $\overline{m}$ upper bound, and often matches exactly $\overline{m}$ patients. Moreover, we show that when Group and Match matches $\overline{m} - 1$ triples, no more triples can be matched among the essential types. This shows that Group and Match is an optimal matching algorithm for the essential types. It also uses entirely 2- and 3-way exchanges.

**Proposition 2** In the absence of any exchange-size constraints, an optimal matching within $E_E$ exactly matches $\overline{m}$ or $\overline{m} - 1$ patients and moreover, Subalgorithm 2, Group and Match, finds such a matching in this subpool using only 2- and 3-way exchanges.

**Proof of Proposition 2:** First observe that by construction Group and Match conducts only 2- and 3-way exchanges. For notational and expositional simplicity suppose $E = E_E$, i.e., $E_E$ is the whole pool. Thus, we drop the argument $E_E$ from $D_X$ and $P_X$ throughout.

**Case 1.** $\lbrack \overline{s}_A, \overline{s}_A \rbrack \cap \lbrack \overline{s}_B, \overline{s}_B \rbrack \neq \emptyset$: We will prove that all triples are matched by the subalgorithm, and
that is $\bar{m}$-many. Without loss of generality assume that
\[
\Delta = n(A - A^* - B) - n(B - B^* - A) \geq 0.
\]
Thus, all $B - B^* - A$ triples are matched in 2-way exchanges with $A - A^* - B$ triples in Step 1 of the Match stage of the subalgorithm.

We first show that $\Delta$ is even. This will be used in the proofs for other cases to rule out certain scenarios.
\[
\Delta = \alpha_A + n(A - A - B) - \alpha_B - n(B - B - A)
= \bar{s}_B - \bar{s}_A + n(A - A - B) - n(B - B - A)
= 2(n(B - A - B) + n(A - O - B) - n(B - A - A) - n(B - O - A)),
\]
showing $\Delta$ is even.

Next, we write down the number of $B - A^* - A$ triples needed to match all $A$ blood-type patients remaining in Step 2 and Step 3 of the Match stage:
\[
\frac{\Delta}{2} + n(A - B - B) + n(A - B - O) - \alpha_A
\]
in Step 2
\[
\frac{\Delta}{2} + n(A - B - B) + n(A - B - O) - \alpha_A
\]
in Step 3
\[
\frac{\alpha_A + n(A - A - B) - \alpha_B - n(B - B - A)}{2} + n(A - B - B) + n(A - B - O) - \alpha_A
\]
\[
\frac{-\alpha_B - \alpha_A + \bar{s}_A - n(B - B - A)}{2} = n(B - A - A) + n(B - O - A) - \alpha_B = n(B - A^* - A).
\]
Thus, all $B - A^* - A$ triples are just sufficient to match all remaining $A - A^* - B$ triples in Step 2 and all $A - B^* - B$ triples in Step 3. Hence, all triples, i.e., $P_A + P_B$ of them, are matched through the subalgorithm. Thus, $\bar{m} \leq P_A + P_B \leq \bar{m}$ where the second inequality follows from Lemma 9. Thus, we have $\bar{m} = P_A + P_B$.

Case 2. “$\bar{s}_A > \bar{s}_B$”: First, we show that $\bar{m}_B = P_B$. We have,
\[
P_B = n(B - B - A) + n(B - O - A) + n(B - A - A) \leq \bar{s}_B \leq \bar{s}_B < \bar{s}_A \leq \bar{s}_A \quad \text{and}
\]
\[
P_B \leq n(B - B - A) + n(B - O - A) + n(B - A - A) + \left[ \frac{\bar{s}_A - \bar{s}_B}{2} \right]
= n(A - O - B) + n(A - B - B) + \left[ \frac{n(B - B - A) + n(B - O - A) + n(A - A - B)}{2} \right]
= \left[ \frac{D_B + D_O}{2} \right].
\]
Since, $\bar{m}_B = \min\{P_B, \lfloor \frac{D_B + D_O}{2} \rfloor, \bar{s}_A\}$, we obtain $\bar{m}_B = P_B$.

In the Group stage, all $A - O - B$ triples are grouped with $A - A - B$’s and all $B - O - A$
triples are grouped with $B - A - A$’s. There are two subcases, $\Delta \geq 0$ and $\Delta < 0$:

**Case 2.1.** “$\Delta \geq 0$”: We have

$$\Delta = n(A - A^* - B) - n(B - B^* - A) = n(A - A - B) + n(A - O - B) - n(B - B - A).$$

First, since $\Delta \geq 0$, Exception (I) is not needed. Second, invoking Exception (II) requires $s_B > s_A$, contradicting we are in Case 2. Thus, no exceptions are invoked in Step 1. Then

$$a := \min \left\{ n(B - A - A) + 2 \min \left\{ n(B - A - A) + n(B - O - A), \left\lceil \frac{\Delta}{2} \right\rceil \right\} \right\}$$

in Step 1

$$+ \max \left\{ 0, n(B - A - A) + n(B - O - A) - \left\lceil \frac{\Delta}{2} \right\rceil \right\}$$

in Step 2

$$\min \left\{ n(B - A - A) + 2n(B - O - A) + 2n(B - A - A) \right\}$$

in Step 3

$A$ blood-type patients are matched in the subalgorithm.

We claim that $a = \overline{m}_A$: Recall that $a \leq \overline{m}_A$ by the upper bound by Lemma 9. Also recall that

$$\overline{m}_A = \min \left\{ P_A, \left\lceil \frac{D_A + D_O}{2} \right\rceil, s_B \right\}$$

and

$$P_A = n(A - A - B) + n(A - O - B) + n(A - B - B),$$

$$\left\lceil \frac{D_A + D_O}{2} \right\rceil = n(B - A - A) + n(B - O - A) + \left\lceil \frac{n(A - A - B) + n(A - O - B) + n(B - B - A)}{2} \right\rceil,$$

$$s_B = n(B - B - A) + 2n(B - O - A) + 2n(B - A - A).$$

Consider the following two subcases:

(a) If $n(B - A - A) + n(B - O - A) \geq \left\lceil \frac{\Delta}{2} \right\rceil$, then

$$\overline{m}_A \geq a = n(B - B - A) + \left\lceil \frac{n(A - A - B) + n(A - O - B) - n(B - B - A)}{2} \right\rceil + n(B - A - A) + n(B - O - A)$$

$$= \left\lceil \frac{D_A + D_O}{2} \right\rceil \geq \overline{m}_A.$$

(b) If $n(B - A - A) + n(B - O - A) < \left\lceil \frac{\Delta}{2} \right\rceil$, then

$$\overline{m}_A \geq a = n(B - B - A) + 2n(B - O - A) + 2n(B - A - A) = s_B \geq \overline{m}_A.$$

Hence, in either case, we have $a = \overline{m}_A$.

Next consider $B$ blood-type patients. Observe that all $B$ patients are matched in the subalgorithm. As $\overline{m}_B = P_B$, we have $\overline{m} = \overline{m}_A + \overline{m}_B$ (the maximum possible number by Lemma 9) patients matched.
Case 2.2. “Δ < 0”: Below we will prove that \( \overline{m} \) patients can be matched except when \( \Delta \) is odd and yet Exception (I) cannot be invoked. In this case, we will show that (a) only \( \overline{m}_A \) \( A \) patients and \( \overline{m}_B - 1 \) \( B \) patients can be matched at most, and (b) our subalgorithm matches exactly that many agents. (Recall that Exception (II) is never invoked in Case 2.) Then by Lemma 9, the result will follow. We analyze Case 2.2 in three subcases:

(a) “Δ < 0 is odd and Exception (I)(A) is invoked in Match stage”:

Since \( s_A > \overline{s}_B \), we have \( n(A - B^* - B) - n(B - A^* - A) - \lfloor \frac{-\Delta}{2} \rfloor > 0 \). Moreover, \( n(B - O - A|B - A^* - A) > 0 \).

Thus,

\[
\begin{align*}
b &= n(A - A - B) + n(A - O - B) + (-\Delta + 1) + n(B - A - A) + n(B - O - A) - 1 \\
&\quad \text{in Step 1} + \text{in Step 2} + \text{in Step 3} \\
&= n(B - A - B) + n(B - A - A) + n(B - O - A) \\
&= \overline{m}_B
\end{align*}
\]

\( B \) patients are matched. On the other hand, the number of \( A \) patients matched is

\[
\begin{align*}
a &= n(A - A - B) + n(A - O - B) + \frac{-\Delta + 1}{2} + n(B - A - A) + n(B - O - A) - 1 \\
&\quad \text{in Step 1} + \text{in Step 2} + \text{in Step 3} \\
&= n(B - A - A) + n(B - O - A) + n(A - A - B) + n(A - O - B) + \left\lfloor \frac{-\Delta}{2} \right\rfloor \\
&= n(B - A - A) + n(B - O - A) + \left\lceil \frac{n(A - A - B) + n(A - O - B) + n(B - B - A)}{2} \right\rceil \\
&= \left\lceil \frac{D_A + D_O}{2} \right\rceil.
\end{align*}
\]

Recall that \( a \leq \overline{m}_A \leq \left\lfloor \frac{D_A + D_O}{2} \right\rfloor \), where the first inequality follows from Lemma 9. Thus, \( a = \overline{m}_A \).

(b) “Δ < 0 is odd and Exception (I)(B) is invoked in Match stage”:

Since \( s_A > \overline{s}_B \), \( n(A - B^* - B) - n(B - A^* - A) - \lfloor \frac{-\Delta}{2} \rfloor > 0 \). Moreover, \( n(B - O - A|B - A^* - A) = 0 \), and at least one exchange can be conducted in Step 1.

Now, the last \( A - A^* - B & B - B^* - A \) 2-way exchange in Step 1 is not conducted: Hence, exactly one \( A - A^* - B \) triple and an even number of \( B - B^* - A \) triples remain unmatched in Step 1. In Step 2, all remaining \( B - B^* - A \) triples are matched with \( A - B^* - B \) triples. Moreover, in Step 3 all \( B - A^* - A \) triples are matched. Thus, \( \overline{m}_B = P_B \) \( B \) patients are
matched. On the other hand,

$$a := \frac{n(A - A - B) + n(A - O - B) - 1 + \frac{-\Delta + 1}{2} + n(B - A - A) + n(B - O - A)}{2}$$

$$= \left\lfloor \frac{D_A + D_O}{2} \right\rfloor$$

A patients are matched (where the equality follows from Equation 12). Thus, $a = \bar{m}_A$.

(c) “$\Delta < 0$ is odd and yet none of the exceptions are invoked, or $\Delta < 0$ is even”:

As $\bar{\Delta}_A > \bar{\Delta}_B$, in Steps 1 and 2 of the Match stage, if $\Delta$ is odd, only one of the $B - B - A$ triples is unmatched, and otherwise, all $B - B - A$ triples are matched. In Step 3, all $B - A - A$ and $B - O - A$ triples are matched with $A - B - B$ triples in 2-way exchanges. Hence, all $B$ blood-type patients, but at most one, are matched. Next we prove the following claim:

Claim: In Case 2.2(c), the subalgorithm matches the maximum possible number of $B$ patients that can be matched, that is $\bar{m}_B - 1\{\Delta$ is odd}.\(^{33}\)

Proof of the Claim: If $P_B$-many $B$ patients are matched, then we are done. Suppose the subalgorithm matches $P_B - 1$-many $B$ blood-type patients. In this case $\bar{m}_B = P_B$. Moreover, $\Delta$ is odd. If we could use all $B$ blood-type patients in exchange, we can collectively provide

at most $\bar{\tau}_B = n(B - B - A) + 2n(B - A - A) + 2n(B - O - A)$ donors to $A$ blood-type patients. Therefore, the maximum number of $A$ patients that can be matched (if it were possible) is: All $A - A - B$’s and all $A - O - B$’s each of which demands one $A$ donor from outside (since $\Delta < 0$, i.e., $n(A - A - B) + n(A - O - B) < n(B - B - A)$ this is feasible), and

$\bar{\tau}_A := \left\lfloor \frac{n(B - B - A) + 2n(B - A - A) + 2n(B - O - A) - n(A - A - B) - n(A - O - B)}{2} \right\rfloor$-many $A - B - B$’s, each of which demands two outside donors. Observe that $\bar{\tau}_A = n(B - A - A) + n(B - O - A) + \left\lfloor \frac{-\Delta}{2} \right\rfloor$. Since $\Delta$ is odd, one of the $A$ blood-type donors provided by one of the $B$ blood-type patients is not used in this upper bound, even though some $A$ patients remain unmatched. Thus, at least one $B$ patient will not be matched in any matching. Thus, the subalgorithm is matching the maximum possible number of $B$ blood-type patients. \(\Box\)

The number of $A$ blood-type patients matched by the subalgorithm is

$$a := \frac{n(A - A - B) + n(A - O - B) - 1 + \frac{-\Delta}{2} + n(B - A - A) + n(B - O - A)}{2}$$

$$= \left\lfloor \frac{D_A + D_O}{2} \right\rfloor$$

by Equation 12. Since we have $a \leq \bar{m}_A \leq \left\lfloor \frac{D_A + D_O}{2} \right\rfloor$, we get $a = \bar{m}_A$.

\(^{33}\)Function $1\{S\}$ gets value 1 when statement $S$ is true and 0 otherwise.
Case 3. \( \bar{s}_B > \bar{s}_A \)`: It is the symmetric version of Case 2 switching the roles of \( A \) and \( B \).

Note that, in the Group and Match subalgorithm, whenever we can, we prioritized \( A - O - B \) and \( B - O - A \) triples in their group while matching. There is a reason for that. Next, we prove that Group and Match not only finds an optimal matching within \( \mathcal{E}_g \), but also matches the maximum possible number of \( A - O - B \) and \( B - O - A \) triples.

**Proposition 3** Consider \( \mathcal{E}_g \), i.e., the subpool with types in \( \mathbb{E} \). Subalgorithm 2, Group and Match, matches the maximum number of \( A - O - B \) and \( B - O - A \) triples possible in any matching; and these numbers are \( \min\{n(A - O - B), \bar{s}_B\} \) and \( \min\{n(B - O - A), \bar{s}_A\} \), respectively.

**Proof of Proposition 3:** First, we show that Group and Match subalgorithm matches \( \min\{n(A - O - B), \bar{s}_B\} \) - many \( A - O - B \) triples and \( \min\{n(B - O - A), \bar{s}_A\} \) - many \( B - O - A \) triples. We prove this for \( A - O - B \)'s (the proof for \( B - O - A \)'s is symmetric). Define \( \kappa \) as the number of \( A - O - B \) triples matched in the algorithm.

**Case 1.** \( \left[ \bar{s}_A, \bar{s}_A \right] \cap \left[ \bar{s}_B, \bar{s}_B \right] \neq \emptyset \)`: All triples in \( \mathcal{E}_g \) are matched by the subalgorithm (by the proof of Proposition 2). Hence \( n(A - O - B) \)-many \( A - O - B \) triples are matched. We have \( \overline{m}_A \)-many \( A \) blood-type patients are matched by Lemma 9. Since \( n(A - O - B) \leq \overline{m}_A \leq \bar{s}_B \), \( \kappa \)-many \( A - O - B \) triples are matched. Thus, \( \kappa = n(A - O - B) \leq P_A = m_A \leq \bar{s}_B \).

**Case 2.** \( \bar{s}_A > \bar{s}_B \)`: In the subalgorithm, triples with \( O \) donors are matched before any other triple in their respective group. Observe that after Steps 1 and 2, either all \( A - O - B \)'s are matched, or no \( A - A - B \)'s are matched, as they are always processed before \( A - A - B \)'s. Similarly, after Step 3, either all \( A - O - B \)'s are matched, or no \( A - B - B \)'s are matched (as remaining \( A - O - B \) types are moved to the \( A - B^* - B \) group after Step 2). Suppose some \( A - O - B \)'s are unmatched. Since we have \( \overline{m}_A \) \( A \) patients matched in this case by the Proof of Proposition 2, all of them are from triples of type \( A - O - B \). Since \( \overline{m}_A \) is the upper-bound of \( A \) patients matched, then maximum possible number of \( A - O - B \) triples are matched. Thus, \( \kappa = \min\{\overline{m}_A, n(A - O - B)\} = \min\{n(A - O - B), \bar{s}_B, \left\lfloor \frac{D_A + D_O}{2} \right\rfloor \} \). Moreover, we have

\[
\left\lfloor \frac{D_A + D_O}{2} \right\rfloor = n(B - A - A) + n(B - O - A) + \left\lfloor \frac{n(A - A - B) + n(A - O - B) + n(B - B - A)}{2} \right\rfloor \\
= \left\lfloor \frac{n(A - A - B) + \bar{s}_B + n(A - O - B)}{2} \right\rfloor \\
\geq \min\{n(A - O - B), \bar{s}_B\}.
\]

Thus, \( \kappa = \min\{n(A - O - B), \bar{s}_B\} \).
Case 3. “$s_B > s_A$”: All $A - O - B$’s are in the $A - B^* - B$ group. If $\Delta \leq 0$ or $\Delta > 0$ is even, then all $A - O - B$ triples are matched following the proof of Proposition 2. If Exception (II)(A) is invoked, then one $A - O - B$ is moved to the $A - A^* - B$ group after Step 1, but is immediately matched in Step 2. Moreover all remaining $A - O - B$’s are matched in Step 3. If Exception (II)(B) or if $\Delta > 0$ is odd and yet no Exception is invoked, then there are no $A - O - B$ triples. Thus $\kappa = n(A - O - B) \leq m_A \leq s_B$.

Hence, $\kappa = \min\{n(A - O - B), s_B\}$. By Lemma 9, this is the maximum number of $A - O - B$ type triples that can be matched (i.e., if there were no other $A$ patients, we would have $m_A = \kappa$).

Theorem 4 Suppose that the dual-donor exchange pool $E$ satisfies the long-run assumption. In the absence of exchange-size constraints, an optimal matching can be found through Algorithm 3, which uses only 2–6-way exchanges. Moreover, the number of patients matched in an optimal matching is given by

$$m - I + \min\{n(A - O - B), s_B\} + \min\{n(B - O - A), s_A\};$$

where $I \in \{0, 1\}$, $s_X$ for $X \in \{A, B\}$ is defined as in Equation 9, and $m$ is defined in Equation system 11.

Proof of Theorem 4: By Proposition 2, $m - I$ patients from the essential triple types $E$ are matched through the Group and Match subalgorithm (in the first step of the sequential matching algorithm in absence of exchange-size constraints) and by Proposition 3, this algorithm also matches the maximum possible number of $A - O - B$ and $B - O - A$ triples. Let $\mu$ be the outcome of this subalgorithm, which is optimal for triples from $E$. By Lemma 8, we can add additionally one triple from types not in $E$ for each $A - O - B$ and $B - O - A$ triple matched in $\mu$. This is the maximum number of triples we can match from types not in $E$ in any matching by the same lemma. Since the number of $A - O - B$ and $B - O - A$ triples matched in $\mu$ is $\min\{n(A - O - B), s_B\} + \min\{n(B - O - A), s_A\}$ (by Proposition 3) then the sequential matching algorithm in absence of exchange-size constraints matches a total of $m - I + \min\{n(A - O - B), s_B\} + \min\{n(B - O - A), s_A\}$ triples, and its outcome is optimal. Matching $\mu$ has exchanges no larger than 3-ways. Since at most one additional triple is inserted in each exchange for each triple matched in the second step of the algorithm, then the final outcome has exchanges no larger than 6-ways.

Theorem 4 implies Theorem 3.