Strategies for Increasing Knowledge, Communication, and Access to Living Donor Transplantation: an Evidence Review to Inform Patient Education

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Abstract

Purpose of review—Inadequate knowledge of the benefits, risks and opportunities for living donation is an important, potentially modifiable barrier to living donor transplantation. We assessed the current state of the evidence regarding strategies to increase knowledge, communication and access to living donor transplantation, as reported in peer-reviewed medical literature.

Recent Findings—Nineteen studies were reviewed, categorized as programs evaluated in randomized controlled trials (8 studies) and programs supported by observational (non-randomized) studies (11 studies). Content extraction demonstrated that comprehensive education
about living donation and living donor transplantation involves multiple learners – the transplant candidate, potential living donors, and social support networks – and requires communicating complex information about the risks and benefits of donation, transplantation and alternative therapies to these different audiences. Transplant centers can help transplant patients learn about living donor transplantation through a variety of formats and modalities, including center-based, home-based and remote technology-based education, outreach to dialysis centers, and social media. Evaluation of these strategies and program themes informed a new Organ Procurement and Transplantation Network (OPTN)/United Network for Organ Sharing (UNOS) public education brochure.

Summary—Increasing transplant candidate knowledge and comfort in talking about living donation and transplantation can reduce educational barriers to pursuit of living donor transplants. Ongoing efforts are needed to develop, refine and disseminate educational programs to help improve transplant access for more patients in need of organ donors.

Keywords
Decision making; health education; living organ donation; living donor transplantation; patient education

INTRODUCTION
Living donor transplantation can provide excellent patient outcomes. The growing organ shortage translates to high morbidity and mortality rates on the deceased donor transplant waiting lists. Living donor kidney transplantation (LDKT) offers patients with end-stage renal disease (ESRD) the best chance for long-term dialysis-free survival. The benefits include faster access to transplant, with less time on dialysis and its associated risks, and superior long-term patient survival compared to both dialysis and deceased donor kidney transplantation (DDKT), at lower cost to the healthcare system. Despite these benefits and a rise in the kidney transplant waiting list to 100,000 persons, LDKT is the least common treatment option, currently performed in 5,800 cases in the United States per year compared to more than 14,000 DDKT in 2017. Living donor liver transplantation (LDLT) can also provide excellent recipient outcomes and reduce morbidity and mortality for patients with liver failure. Potential benefits of LDLT over deceased donor liver transplantation include the ability to control the timing of the transplant before the patient becomes too ill for transplant. The liver transplant waiting list currently comprises 14,000 persons. LDLT is performed for approximately 350 U.S. patients each year, compared to 7,700 deceased donor liver transplants in 2017.

Based on data and surveys supporting positive public perceptions of living donation, a number of initiatives are underway to increase awareness of living donor transplantation. One example is an American Society of Transplantation Consensus Statement promoting LDKT as the “best treatment option” for eligible patients with kidney failure. The 2017 Kidney Disease: Improving Global Outcomes (KDIGO) “Clinical Practice Guideline on the Evaluation and Care of Living Kidney Donors” recognizes that public awareness of opportunities for living donation should be improved through education. While the need and benefits are clearly established, multiple barriers to living donor transplantation exist.
Barriers include inadequate education and outreach, the complexity and potential inefficiencies in the living donor evaluation process, disincentives such as uncompensated costs for the donor, and ongoing needs to support and advance donor safety and risk assessment to sustain public trust in the donation process. Each barrier is a potential target for focused interventions (Figure 1).6

Educational barriers to the pursuit of potential living donors by transplant candidates, or more specifically, inadequate knowledge of the benefits, risks and opportunities for the procedure, are important and modifiable. Comprehensive education about living donation and living donor transplantation involves multiple learners – the transplant candidate, potential living donors, and social support networks – and requires communicating complex information about the risks and benefits of living donation, transplantation and alternative therapies to these different audiences.7 Transplant programs can help transplant candidates learn about living donor transplantation through a variety of formats and modalities, including center-based, home-based and remote technology-based education, outreach to dialysis centers, and social media. Understanding the impact of such interventions is necessary to inform center choice and investment in programs suitable for their resources, processes, and patient populations, and to encourage patients to seek and participate in available programs at their centers.

To reduce educational barriers to living donor transplantation, a work group of the Organ Procurement and Transplantation Network (OPTN)/United Network for Organ Sharing (UNOS) Living Donor Committee formulated a project to create a new educational brochure for transplant candidates. As a foundational step in the brochure development, the work group reviewed the state of current evidence on strategies to increase knowledge, communication and access to living donor transplantation, as reported in peer-reviewed medical literature, as well as some emerging programs that have not yet been formally evaluated. This article describes the processes and findings of our evidence review.

EVIDENCE REVIEW METHODOLOGY

Process of Educational Program Identification

The work group identified and reviewed studies of living organ donor educational programs reported in the medical literature. We queried the MEDLINE electronic database for reports published through February 28, 2017 using the Medical Subject Headings (MeSH) terms “living donor* or living organ donor*”, “live donor* or live organ donor,” “live donor kidney transplantation”, “live donor liver transplantation”, “living donation”, “tissue donor*”, “kidney transplant*”, “liver transplant*”, “education”, “patient education,” “health education,” “education program,” “education session,” “teaching or teaching session,” “counseling,” “decision aid,” “pamphlet,” “handout or patient education handout,” “audiovisual aid” “teaching materials,” “consumer health information,” “health knowledge, attitudes, practice,” and “healthcare disparities”. Searches were limited to articles published.
in English. Manual review of the reference lists of identified articles was also performed. Each study was reviewed to extract a brief program description (Tables 1 and 2), a description of supporting evidence, and information about study limitations or other considerations. One work group member performed the primary extraction for each identified article, and a second work group member confirmed the extraction.

The project was not designed as a systematic review in which only a narrow range of studies are assessed. Rather, the work group sought to identify the full range of living donor transplant and donation educational programs that have been developed and evaluated. Studies were categorized by methodologic rigor based on study design, ranging from randomized controlled trials, to observational or non-randomized studies. Emerging programs that have not yet been formally evaluated or that do not yet have published data were described but were not included in the data extraction.

Based on the available evidence, the work group identified key elements of effective educational strategies. The evidence is predominantly related to LDKT. While educational programs focused on LDLT are also critically important, there is currently a paucity of evidence for the effectiveness of educational programs related to this transplant option.

EDUCATIONAL PROGRAM SUMMARIES AND EVIDENCE

We identified 19 studies meeting selection criteria. The identified educational programs are described below, categorized as programs evaluated in randomized controlled trials (8 studies) and programs supported by observational (non-randomized) studies (11 studies).

Each educational program description includes a brief summary of: 1) the program’s overall approach (further programmatic information is presented in Tables 1 and 2); 2) the evidence supporting the program’s efficacy; and 3) other commentary. Following the program summaries and evidence, we briefly describe several emerging programs that have not yet been formally evaluated. Finally, we summarize some of the educational programs’ commonalities with respect to design, content and delivery teams and suggest next steps to advance patient education.

Programs Evaluated in Randomized Controlled Trials (Table 1)

While randomized controlled trials offer the highest standard of evidence, only a limited number of trials have been performed to evaluate educational interventions related to living donor transplantation. Most included knowledge and attitudes as endpoints, while only a subset reported living donor evaluations or access to living donor transplantation.

**“House Calls” or Home-Based Education Program**

**Program:** “House Calls” is a home-based (HB) intervention in which trained transplant health educators perform a 60 to 90-minute LDKT education session involving family and members of the candidate’s social network. [Figure 2]

**Evidence:** In one single-center trial, 152 kidney transplant candidates were randomized to clinic-based (CB) education or CB+HB. Compared to CB alone, significantly more patients...
who received CB+HB had living donor inquiries (83% vs 64%), donor evaluations (60% vs 35%) and LDKT (52% vs 30%) one-year following the intervention. Both groups demonstrated increased LDKT knowledge after CB education, but HB education led to the largest increase in LDKT knowledge, and also to greater willingness to discuss LDKT with others and a decrease in LDKT concerns.

A secondary analysis provided details on culturally sensitive aspects of the intervention: This analysis found that increases in both donor evaluations and LDKT at 1 year with CB +HB compared to CB were larger for black (versus white) candidates. Notably, the LDKT rate for black patients who received HB education mirrored the LDKT rate of white patients before the intervention; i.e., the intervention appeared to eliminate the initial racial disparity in LDKT at that center.

In a second single-center trial, 152 black kidney transplant candidates were randomized to the HB intervention, CB education in a group session with other patients and their guests at the transplant center, or CB individual patient counseling. Although rates of LDKT did not differ significantly between the three groups at the 2-year endpoint, patients receiving HB education were more likely than patients in the other two groups to have at least one donor inquiry (82% vs. 61% vs. 47%) and evaluation (65% vs. 39% vs. 27%). Patients in the HB group also had higher LDKT knowledge, fewer LDKT concerns, and higher willingness to talk to others about donation 6 weeks after intervention.

Commentary. Results of these two trials may be limited by possible self-selection bias among those who gave consent and completed the study (i.e., such patients already may have been thinking about LDKT). In the first trial, there was a higher dropout rate among patients randomized to receive HB education, suggesting further self-selection, and the drop-out rate was higher among black (versus white) patients. The degree to which trial results generalize to other groups of patients and to other regions of the United States is unknown. Training and supporting staff to perform the HB intervention may be too costly or logistically prohibitive for some centers.

“Kidney Team at Home” Home-Based Family Intervention

Program: This program is a Dutch adaptation of the “House Calls” intervention.

Evidence: In a single-center trial, 163 patients referred for transplant evaluation or on the DDKT waiting list were randomized to usual education or usual care plus + HB education. All participants had been unable to find a potential living donor prior to enrollment. Patients in the CB+HB group showed significantly more improvements in LDKT knowledge and communication compared to the control group. Participating members of the patient’s social network also showed significant increases in knowledge and positive attitudes related to discussing kidney replacement therapies and willingness to donate a kidney, and a decrease in risk perception. Excluding patients who died or who received a DDKT, those who received CB+HB education (versus CB education alone) had significantly more living donor inquiries (32% vs. 74%), evaluations (17% vs. 64%) and actual LDKTs (10% vs. 44%) at 6 months following the intervention. In a second trial involving patients who had not yet
initiated renal replacement therapy, patients who received CB+HB education (versus CB education alone) were substantially more likely to receive a preemptive LDKT (65% vs 39% of those who received LDKT within 2 years of intervention).\textsuperscript{12}

Commentary. The follow-up period for LDKT in the first trial was relatively brief and the trial results may be limited by possible self-selection bias among those who gave consent and completed the study. Of note, the LDKT rate at the study center already was very high (~70%), so patients enrolled in the trial may have faced more substantial barriers to LDKT than other patients at the center, given the baseline success at this center. Drop-out was associated with limited social networks, suggesting that the intervention may be less effective in patients with fewer family members and friends. The second trial well-received by those who used it, but had a high non-participation rate among the target population of 60%, with many declining for the very barriers that such educational programs aim to overcome (concern for burdening others; concern for insufficient social network). Also, although aimed at encouraging pre-emptive LDKT, a large proportion of participants did not progress to needing renal replacement during the 2-year study period, demonstrating the challenge of appropriately timing transplant education for pre-dialysis patients.

“Talking About Live Kidney Donation” (TALK) and “TALK-Social Worker” Intervention programs

Program: The TALK education intervention consists of a video and booklet on LDKT,\textsuperscript{13} The “TALK-Social Worker” intervention combines the video/booklet with two hour-long sessions delivered by a social worker either in the patient’s home or in the transplant center.\textsuperscript{14} In the first session, the social worker distributes the video and booklet and helps the patient identify barriers to considering or pursuing LDKT. The patient is asked to invite family/friends to a second session. At the second session, the social worker helps social network members identify barriers to discussing LDKT and living donation. Problem-solving and motivational interviewing techniques are employed at both sessions. [Figure 3]

Evidence: In this trial, 130 patients were randomized by site (nephrology practice) into three study arms (usual care with nephrologists, and one of the two TALK conditions). Eligible patients were drawn from academically affiliated or community nephrology practices, and were not yet on dialysis (although they may have had a transplant evaluation). The primary outcome was whether patients engaged in any one of five behaviors reflecting discussion and/or pursuit of living donor transplantation (discussions with family, discussions with physicians, initiating transplant evaluation, completing transplant evaluation, identifying a potential living donor). Engagement in at least one behavior over 6 months was 30% with usual care, 42% with TALK-education, and 58% with the “TALK-Social Worker” intervention. Results were not significantly different between the two TALK arms. The most common behaviors were talking to family and physicians. No differences were observed between groups in interest or concerns about LDKT.

Commentary. The follow-up period was relatively brief, and the study did not consider actual receipt of LDKT. The educational materials did not provide
extensive information on risks and benefits of LDKT, and financial barriers were not considered in the TALK interventions. Only 67% of patients in the “TALK-Social Worker” intervention attended both sessions, and 19% of patients attending the second session did not bring family/friends. More complete uptake of the interventions may have been limited by the high percentage of pre-dialysis patients who may not have been “primed” or ready for focused discussion of LDKT and living donation.

“Living ACTS” (About Choices in Transplantation and Sharing) [Figure 4]

**Program:** “Living ACTS” (About Choices in Transplant and Sharing) was designed as a culturally sensitive intervention targeted to African-American patients. The intervention consists of a video and booklet.

**Evidence:** In a single-center trial, 296 black adults who presented for transplant evaluation were randomized to receive standard education plus the “Living ACTS” intervention, or standard transplant education plus an attention-control video about exercise. Standard education comprised a web-based course (transplant-specific education) completed before the evaluation. Knowledge about LDKT and willingness to talk with family about living donation were assessed before randomization, immediately after education, and 6 months later. Subjects in the “Living ACTS” group showed a significantly greater increase in knowledge about living donor transplantation immediately after the intervention than did controls; this was maintained through the 6-month follow-up period. Although the intervention group showed significantly higher willingness to talk with family about living donation, this difference was evident at baseline and did not change over time. Intervention participants endorsed greater perceived benefits of living donor transplantation than did controls immediately after the intervention, but this effect did not persist at the 6-month follow-up.

**Commentary:** Although 96% of patients approached agreed to enroll, the sample was one of convenience and generalizability is not known. In addition, the analyses did not follow an intent-to-treat approach and included only those with complete follow-up data. Because no information was collected from patients on educational activities or interactions with family regarding living donation after they returned home, it is difficult to speculate on why the intervention affected only patients’ knowledge.

“Infórmate” Culturally Targeted Website for Hispanic Transplant Candidates

**Program:** The authors developed a bilingual, culturally targeted website, “Infórmate”, for Hispanic transplant candidates and members of their social network. The website was designed to overcome many of the known sociocultural barriers to LDKT among Hispanics, including misconceptions about living donation, language barriers, immigrant status issues, financial concerns, and distrust of the healthcare system.

**Evidence:** In a two-center trial, 282 new kidney transplant candidates were randomized to receive access to the website or to usual care. Usual care comprised attending a transplant
education class on the day of their initial evaluation. Patients and their family members in both groups received this standard care; however, one group was also given access to “Infórmate” before attending the class. Patients and family members in both groups had significant improvements in LDKT knowledge from pre-intervention to the 3-week follow-up; however, knowledge gains were larger for patients assigned to website group. [Figure 5]

Commentary. The majority of study participants were family members of the transplant candidate. Knowledge gains were not presented by participant type, thus it is unknown whether knowledge gains occurred equally for patients and family members. There is also a possibility of intervention contamination, as the website intervention occurred before a group education session including all study participants. Patients and family members who had access to the website may have shared its content and information with participants in the usual care group, thus potentially suppressing the effects of the website intervention. Finally, the study evaluated changes in knowledge only, so it is not known whether the website impacts rates of living kidney donation or LDKT.

Written Educational Materials followed by Structured Educational Session

Program: In this intervention, patients newly referred for transplant evaluation are mailed written materials about living kidney donation and the benefits of LDKT. Approximately two weeks later, patients and their invited guests attend a 2-hour educational session at the transplant center.

Evidence: In a single-center trial, 100 patients who presented for their initial transplant evaluation were randomized to receive either usual care or usual care plus the enhanced LDKT educational program. Usual care comprised informal discussions with providers about all aspects of kidney transplantation as patients progressed through the evaluation process. The primary outcome was potential living donor contacts with the transplant centers within 3 months after the intervention. Also, patients completed questionnaires about LDKT knowledge and ESRD treatment preferences before and 2 weeks after the intervention. The two groups did not differ significantly in the number of potential living donors who contacted the center. Extending the follow-up period to 6 months also did not yield significant between-groups differences. Patients in the intervention group who had not identified LDKT as their treatment preference initially were more likely to state that this was now their treatment preference 2 weeks after the intervention.

Commentary. The trial only enrolled 50% of the intended sample size, and the authors acknowledge that the study was under-powered. Although LDKT knowledge was assessed, these results are not reported. Details are lacking regarding the content of usual care, making it difficult to determine the nature and dose of any LDKT information the control group may have received. Finally, the drop-out rate for the intervention was very high (30% did not attend the education session), suggesting further self-selection.
Programs Supported by Observational (Non-Randomized) Studies (Table 2)

A variety of additional programs have been developed that may be offered at transplant centers or elsewhere but that have not yet been tested in randomized controlled trials. Thus, efficacy in improving knowledge or identification of potential living donors has not been fully established. These programs relied on quasi-experimental designs (e.g., use of historical controls or comparison groups), or single group pre-post observational designs, often in small pilot samples. Despite the early nature of the evidence, the programs below are noteworthy for their creativity in developing strategies using peer-mentors, social media tools, and mobile decision aids.

“Patient Navigators” in Community-Based Nephrology Practices

Program: Social workers, or “Patient Navigators”, are placed in community-based nephrology practices after receiving training in transplantation, living donation, and cultural sensitivity from transplant physicians, nurse coordinators, and social workers. The social workers meet with patients before or after their appointment with the nephrologist and, upon request, schedule additional individual appointments with patients.

Evidence: In a non-randomized observational study conducted over 58-months, patients referred to the transplant center from two nephrology offices with a social worker-navigator were compared to patients referred from 90 nephrology offices without patient navigation. The investigators retrieved outcomes data on 2,722 transplant candidates and 424 transplant recipients at their center. Social worker-navigators had a mean of 6 contacts per patient (range: 1 to 32). Patients from nephrology practices with a social worker-navigator were more likely to have potential living donor inquiries (38% versus 34%) and initial screening (30% versus 26%) compared to patients from nephrology practices without navigators. However, the two groups did not differ significantly in rates of complete living donor evaluations or LDKT (4% in both groups).

Commentary. The “dose” of the intervention was not standardized, which may be relevant to efficacy of the program. Patients who had more contacts with social worker-navigators may have progressed more rapidly through the transplant evaluation process and had more interest in LDKT. Social workers were hired as full-time employees, which may make the intervention cost-prohibitive for many transplant programs to adopt. Finally, there was no defined, consistent time period from navigation to the observed outcomes (living donor inquiries and evaluations, LDKTs) across the sample.

“Living Donor Family Education” Program

Program: Designed and then refined across a series of publications, this program is offered at the transplant center during the evaluation process to “recipient families,” which includes relatives as well as friends of candidates for kidney or simultaneous kidney-pancreas transplant. The intervention comprises a didactic presentation and discussion led by nurse coordinators familiar with living donation, LDKT, and related literature. The intervention also includes an educational video.
Evidence: In an initial publication, after the program had been offered for over one year, the percentage of patients who had potential living donors submit blood samples to screen for candidacy in the group receiving the new program were compared to the percentage of patients screened before the program existed. Among 1,363 patients registered on the transplant waiting list during the 54-month study period, a significantly higher percentage of patients had at least one potential living donor tissue typed after versus before the program was implemented (39% vs. 33%). The program was particularly effective for blacks and older patients, who had higher rates of living donor screening after versus before program implementation (36% vs. 28% for blacks; 31% vs. 19% for older patients). In additional analyses, the authors observed that the rate of kidney transplant was significantly higher after the program’s initiation than before, but post-transplant survival rates were not affected. In subsequent analyses over a 4-year period after program inception, the authors continued to find increased rates of potential living donors submitting blood tests after versus before program implementation, but findings were inconsistent in terms of subgroup effects.

Commentary: The results are primarily limited by reliance on a historical comparison group; rates of potential living donors submitting blood tests may have increased naturally over time even without the intervention. In addition, although the authors conclude that it is important to have ethnically appropriate teaching materials, they did not directly evaluate whether that specific component contributed to any effects. The study included only a single center and generalizability is not known.

“Seminars on Saturdays” Program

Program: The intervention is targeted toward (but not exclusive to) preemptive kidney transplantation, and consists of formal talks, facilitated discussion during break-out sessions, audiovisual aides, and an information packet, along with follow-up contact by a transplant coordinator to determine if more information is needed. The program is offered by a multidisciplinary team every 2 months (6 times per year) on a Saturday to pre-dialysis patients and their family/friends, as well as to all patients on the kidney waitlist, and to all dialysis patients not yet waitlisted but deemed suitable for transplant. [Figure 6]

Evidence: Analyses are based on comparisons of outcomes collected annually during a period spanning 3 years before and 3 years after the program began to be offered at a single center. The investigators report a growth in program attendance and a significant increase in the percentage of LDKTs that were preemptive at their center (from 28% to 44%). The percentage of non-blood-related donors among total living donors increased from 28% to 41% (statistical significance not reported). Preemptive LDKT that were from non-blood-related donors significantly increased (from 25% to 58%). The authors also found that the percentages of minority donors increased from 39% in earlier years to 52% in later years (statistical significance not reported).

Commentary: The results focus on comparisons of annual data and do not directly compare outcomes in individuals who attended an intervention session versus those who did not attend. The changes over time cannot be specifically attributed to launching the intervention. The analyses do not appear to specifically analyze
trends over time but instead compare rates across 6 nominal groups (defined by year). In other words, the authors report a difference across the 6 groups defined by year but do not test directly that the percentages monotonically increase by year. Some effects are not subjected to any statistical test.

“Live Donor Champion” Program

**Program:** The “Live Donor Champion” (LDC) program was designed on the premise that training a “champion” or advocate to share about the transplant candidate’s disease and need for an organ donor would help identify more living donors. In the LDC program, each transplant candidate identifies one person to be their “Live Donor Champion”, who is then trained to talk about kidney disease, organ donation and transplant, and skills for initiating conversations. Anti-coercion training is an important part of the curriculum. Champions are provided with educational materials, “business cards” with contact information, and other resources from the transplant center.

**Evidence:** Pilot data come from a single-center prospective cohort study of 15 adult kidney transplant candidates who had been on the DDKT waitlist for at least 3 months and had no potential living donors at the time of enrollment. Each candidate identified someone from their social network to serve as their LDC. LDC comfort in initiating conversations about transplantation increased over time. Twenty-five potential donors contacted the center on behalf of LDC participants; four participants achieved LDKT and three additional participants had donors in evaluation, compared to none of the 15 matched controls. [Figure 7]

**Commentary:** This was a non-randomized, small pilot study subject to selection bias because those patients who nominated “LDCs” may have been more motivated and willing to pursue LDKT. While nearly half of the “LDC” were black, it is unknown whether the intervention yielded the same benefit regardless of race. The lengthy duration of the intervention (6 months) is a practical limitation, especially for those patients seeking preemptive LDKT. Finally, patients with smaller social networks may benefit less from the intervention, particularly if they are unable to identify a champion.

“Hispanic Transplant Education” Program

**Program:** This intervention is designed to be a linguistically and culturally competent educational program for Hispanic transplant candidates, using an established framework for standardizing the measurement and reporting of high quality, culturally competent care. The intervention consists of two educational sessions offered at the time of patients’ evaluation for transplant at the center.

**Evidence:** The study used a pre-post design in which 43 patients and 70 family members completed pre-intervention measures, received the intervention, and then immediately completed post-intervention measures. Family members’ and friends’ attitudes about being a living kidney donor were also assessed. Knowledge level significantly increased among both patients and family/friends. Knowledge was higher at post-test among patients.
than in family/friends, and among younger and more educated individuals. Pre-to-post attitude change was noted in some areas: patients became more motivated to pursue LDKT and felt more favorable about it. Family/friends increased in willingness to become living donors and became more motivated to begin the living donor evaluation. High levels of satisfaction with the intervention were noted.

**Commentary.** The resources needed to mount this type of intervention may not be available to all transplant programs. The study did not include a comparison group and the analyses focused on patients who completed the intervention. Thus, the robustness and generalizability of the findings are unknown. There was no follow-up period to determine if knowledge and attitudes were maintained over time. It is unclear whether improved attitudes regarding living donation and motivation to pursue living donation translated into an increase in the number of living donor inquiries among those family/friends.

### Educational Intervention Based in Peer Experiences

**Program:** This intervention was developed to expose liver transplant candidates (and potential living liver donors in their social network) to the experiences of former living liver donors. The intervention includes a booklet and video (available in 6 languages), and a website that includes the same written and video materials.

**Evidence:** Across 5 transplant centers, the investigators compared 338 liver transplant candidates who received the intervention to 437 patients who received “usual education” at those same centers before the intervention was offered. Primary outcomes were patients’ living donation/transplantation knowledge and feelings of self-efficacy to discuss living donation with others. Patients in the intervention group reported significantly more knowledge and higher levels of self-efficacy than those in the comparison group. Patients in the intervention group were significantly more likely to have talked about living donation with others than those in the comparison group. The total number of family/friends who presented for donor evaluation and the number of actual living donors both increased from before to after intervention over a 3-year period. Among those in the intervention group who were known to either have viewed the intervention materials (27% of the group) or not viewed them (58% of the group), those who viewed them showed significantly higher self-efficacy for discussing living donation with others.

**Commentary.** The results are primarily limited by reliance on a historical comparison group; other elements may have led to the differences observed in knowledge and self-efficacy, and the numbers of donor candidate evaluations and ultimate donations may also have changed over time due to other factors. Only a minority of patients in the intervention group reported viewing the intervention materials, and so the actual exposure to the intervention was limited.

### “Pre-Dialysis Education” Program

**Program:** A “Pre-Dialysis Education” program (PDEP) was developed as a training kit using visual and written information cards and comprising 6 modules targeted towards the degree of kidney disease experienced by patients with chronic kidney failure.
Evidence: The intervention was evaluated in a single university hospital in Turkey in a non-randomized retrospective design. Among 88 LDKT recipients over an 8-year period, 61 patients who received PDEP were compared to 27 patients who received transplants before PDEP was available. Significantly more PDEP patients had preemptive LDKT than those who did not receive PDEP (43% vs. 19%). PDEP participants were more likely to receive preemptive transplants from spouses, siblings and non-parent relatives, and less likely to receive preemptive transplants from parents, than patients not exposed to PDEP.

Commentary. Very limited detail is available regarding the conceptualization, development or specific content of the PDEP modules, challenging both the ability to replicate the intervention in other settings, and the evaluation of strengths and limitations of the intervention.

“Living Donation Education” Program, center-based LDKT education for black and Hispanic patients

Program: The “Living Donation Education” (LDE) program was developed to address common barriers to LDKT and living kidney donation among black and Hispanic patients. The intervention is delivered in two phases at the transplant center. In Phase I, patients and family members attend a formal education session with healthcare providers. In Phase II, patients and family members are invited to meet with a living donor and recipient pair. The study originally was designed to also examine HB education developed by others; however, this group was dropped after only one patient chose to participate in a HB education session.

Evidence: In a single-center study, 294 patients on the kidney transplant waiting list were informed about LDE by telephone and encouraged to attend the program. Of these, 75 patients attended the Phase I educational session and 7 patients returned for the Phase II session. The authors tracked the number of patients who attended the LDE and examined whether they had a higher likelihood of living donor inquiries, evaluations, and transplants compared to patients who did not attend an LDE session but who attended the transplant program’s regular patient support group. There were no statistically significant differences between the two groups on any of these outcomes. Also, the two groups did not differ in changes in LDKT knowledge, willingness, concerns, or readiness.

Commentary. The study is limited by low statistical power, non-randomized design, selection bias, and cross-contamination. Few patients attended the sessions, particularly the second session. It is possible that interest in LDE was limited because the transplant program in which the study occurred was not able to perform LDKT for a year due to program issues that occurred shortly after study initiation. While the program was permitted to evaluate potential living donors, the restriction on performing LDKT could have adversely impacted study participation rates and outcomes. Also, half of the patients receiving the LDE intervention also participated in the control condition by attending one or more support group sessions. The initial study design included a HB education arm which could not be assessed due to lack of participation.
Social Media/Facebook Apps

**Program:** A Facebook smartphone app was developed that enables transplant candidates to share their need for a living donor through social media, providing an indirect means for communication with a broad social network.29

**Evidence:** In a pilot study among 54 adult transplant candidates without potential living donors at the time of enrollment, the majority of candidates using the smartphone app reported it to be “good” to “excellent” with respect to the installation process, readability, simplicity, clarity and content.29 Compared to matched controls, patients using the app were nearly 7-times as likely to have a potential donor come forward on their behalf within 10-months after enrollment. [Figure 8]

**Commentary.** Limitations of this initial study include the small sample size, non-randomized design, and potential selection bias from enrollment of more motivated or technologically sophisticated candidates. Future studies should assess the impact of the Facebook smartphone app on conversion of donor referrals to LDKT, cost-effectiveness (including consideration of staff time for referral intake and screening), and the quality of patient shared-decision making about transplant options and donation.

“My Transplant Coach” Decision Aid

**Program:** A mobile iOS (iPad) app was created utilizing animated patient education and individualized, risk-adjusted outcomes following kidney transplantation to improve overall transplant knowledge in kidney transplant candidates. The patient-centered information allows candidates to consider the relative benefits of LDKT and various types of DDKT (e.g., low-moderate kidney donor profile index (KDPI), high KDPI, or an Increased Infection Risk donor). 30

**Evidence:** In a pilot study of 81 patients at two large US transplant programs, the participants were guided through the app by a research assistant prior to receiving any standard educational sessions at their transplant program. Participants completed pre- and post-surveys of the app to assess comfort with technology, knowledge of transplant options and acceptability and cultural competency of the app. Overall, the app resulted in significant increases in patient knowledge of transplantation, acceptance of LDKT and DDKT options and comfort in discussing transplantation with others.

**Commentary:** A strength of the app is the presentation of patient-centric education based on the individual candidate’s demographic and clinical information and relevant data for their anticipated Donation Service Area. Though the app scored well for overall impression and appropriateness for participant’s race/ethnicity, the study participants were of modestly higher education than the average ESRD population and already possessed moderate health literacy. The app improved transplant knowledge and understanding of outcome differences between LDKT and DDKT options; however, differences were observed based upon pre-existing access to and comfort with technology. The app is also only available in English, and so may not fully address the educational needs of candidates with limited...
English skills; however, a Spanish translation is expected in the next version of the app. The study did not assess the impact of the app on the participants’ actual pursuit and acceptance of organs.

**Emerging Programs**

Other educational programs focused on LDKT and living donation are emerging, including in planned trials, but have not yet been formally evaluated. Examples of such emerging programs include:

- **“Explore Transplant at Home.”** This program will offer video and print educational resources about DDKT and LDKT developed as part of the “Explore Transplant” program to socioeconomically disadvantaged dialysis patients in their homes, with or without supplementary telephone interactions with a health educator. Thus, the study is focused on the impact of home-based delivery of previously developed educational resources.

- **“ELITE” Program.** This program will use and adapt the “Explore Transplant” materials as part of the education of potential adult kidney transplant candidates attending their initial transplant evaluation at one transplant center, in addition to usual educational materials. The program will focus on knowledge of living donation and living donor transplantation, and include a video and meeting with a transplant educator.

- **“Your Path to Transplant.”** This study will evaluate materials from the “Explore Transplant” program in a computer-tailored format. The novel aspect of this program is examination of a validated transplant decision-making measure that can be delivered either in person, via the telephone, or electronically via a computer.

- **“iChoose Kidney.”** “iChoose Kidney” was developed by a multi-disciplinary team as a shared decision aid for clinicians to present individualized risk predictions of short-term patient survival with kidney transplantation versus dialysis to patients. It is available through the Internet (http://ichoosekidney.emory.edu/) or a mobile (iOS) app. “iChoose Kidney” presents the patient survival or mortality risk in a patient-friendly manner using both graphical and numeric representations. A randomized trial to assess knowledge, treatment preferences and access to transplant including LDKT is underway.

**Educational Program Commonalities: Design, Content and Delivery Teams**

While education programs differ in detail, published studies suggest some commonalities among effective programs, including provision of information early in the organ disease process, an opportunity to include family and friends, ideally in a comfortable setting, and opportunities to talk with living donors and living donor transplant recipients (Table 3). Many of the published interventions seek to help transplant candidates identify individuals from their social network to participate with the candidate in educational sessions. A common design theme focuses on supporting productive communication between transplant candidates and members of his or her social network on organ donation and transplantation,
often in the presence of a transplant medical expert to answer questions and concerns for both parties, or after training the patient or an advocate to communicate with the social network. Some of these programs provide educational materials (e.g., brochures, videos, decision aids) prepared for both the transplant candidates and potential donors. Social media tools for sharing a patient’s need for an organ donor safely and effectively with their social network are also emerging.

Regarding content, effective educational programs generally include core subject matter related to the need, risks, benefits and outcomes of living donation and living donor transplantation, compared to other treatment options (Table 4).

To deliver education, effective living donor transplant educational programs include a cross-section of transplant professionals and volunteers (Table 5). Educational programs need not rely disproportionately on any one category of transplant professional. Multi-disciplinary transplant teams, often supported by peers including living donors and transplant recipients, help provide and support effective education.

Transplant center education programs can also positively influence patients’ use of the internet in connection with organ transplantation. First, transplant programs can help vet the factual basis of online information when patients begin to consider their options. This is particularly important given the amount of information – and misinformation – about living donation accessible on the internet. Second, educating patients and their friends and family about living donation can increase their appropriate and constructive communication about living organ donation and transplantation on social media, as described above. Importantly, literacy-related disparities in technology access and use are widespread, such that patients with lower levels of literacy are less likely to own smartphones or to access and use the internet, particularly for health reasons. Thus, future interventions should consider these disparities and ensure that health promotion activities do not further exacerbate disparities.

**Conclusions and Next Steps**

Living donor transplantation is an important treatment option for end-stage organ disease that can be performed safely to benefit many patients and society. However, living donor transplantation has not increased in response to the growing organ shortage. Increasing transplant candidate knowledge and comfort in talking about living donation and transplantation can reduce educational barriers to pursuit of living donor transplants. The likelihood that transplant candidates will identify someone willing to consider living donation can be increased by the provision of clearly stated educational information, involving the patient’s social network, and repeated over phases of care. Electronic-health technology is emerging as an efficient modality for education delivery, but more comparative work is needed to determine whether electronic-learning, telehealth, or telephone-based education programs are as effective as programs that deliver education with an in-person educator. New technology-based tools, such as social media, also show promise as strategies to help patients expand their social network, but require careful development with attention to online privacy and safety, and efforts to avoid disadvantaging those with low access to technology. A new patient-focused educational brochure, “How to Find a Living Donor: Make Your Transplant Happen”, describes some evidence-based education strategies and is
available as a print-on-demand resource from the OPTN website (https://optn.transplant.hrsa.gov/media/2267/find_a_living_donor.pdf).

Unanswered questions related to living donor transplant educational interventions include how much educational content (i.e., “dose”) should be provided and at what time points in the transplant evaluation pathway. It is also unclear whether the dose should vary based on patient characteristics (e.g., disease state, race/ethnicity, size of social network, etc.) and to what extent interventions should be tailored to these patient characteristics. There are known disparities in access to living donor transplant based on race/ethnicity, literacy and health literacy, and other factors, and efforts to improve equity for non-white patients and those with lower levels of socioeconomic status and health literacy are important priorities. Empirical studies of approaches to support and increase living donor transplantation including formal evaluations of education, incorporating assessment of transplant rates as hard outcomes and associated costs, are feasible and necessary to advance evidence-based, effective practices. Robust efforts to develop and evaluate strategies for increasing knowledge, communication and access to living donor transplantation must to continue to support opportunities for more healthy, willing persons to give the gift of life to patients in need.

Acknowledgments

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ABBREVIATIONS and ACRONYMS

<table>
<thead>
<tr>
<th>ACTS</th>
<th>“About Choices in Transplant and Sharing”</th>
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<tr>
<td>CB</td>
<td>Clinic-based</td>
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<tr>
<td>DDKT</td>
<td>Deceased donor kidney transplantation</td>
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<td>DSA</td>
<td>Donation Service Area</td>
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<td>ESRD</td>
<td>End-stage renal disease</td>
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<td>HB</td>
<td>Home-based</td>
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<tr>
<td>KDIGO</td>
<td>Kidney Disease: Improving Global Outcomes</td>
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<td>KDPI</td>
<td>Kidney Donor Profile Index</td>
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<td>LDC</td>
<td>“Live Donor Champion”</td>
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LDE Living donation education
LDKT Live donor kidney transplantation
LDLT Living donor liver transplantation
MeSH Medical Subject Headings
OPTN Organ Procurement and Transplant Network
PDEP “Pre-Dialysis Education” program
TALK “Talk About Live Kidney Donation”
UNOS United Network for Organ Sharing

**APPENDIX. Resources for more information**

<table>
<thead>
<tr>
<th>Program</th>
<th>Investigators</th>
<th>Contact</th>
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<tr>
<td>“House Calls”</td>
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<td></td>
<td>Beth Israel Deaconess Medical Center</td>
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<td></td>
<td>Boston, MA, USA</td>
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<tr>
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<td></td>
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<td></td>
<td>Rotterdam, The Netherlands</td>
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<td>Duke University School of Medicine</td>
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<tr>
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<td></td>
<td>Northwestern University School of Medicine</td>
<td><a href="http://www.informate.org">www.informate.org</a></td>
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<td>Chicago, IL, USA</td>
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<td>Calgary, Alberta, Canada</td>
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<td>“Living Donor Family Education”</td>
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<td>University of California</td>
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<td></td>
<td>Irvine, Orange, CA, USA</td>
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<tr>
<td>“Seminars on Saturdays”</td>
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<td>Hispanic Transplant Education</td>
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<td></td>
<td>New York Center for Liver Transplantation</td>
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<td>“Pre-Dialysis Education”</td>
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<td>Mustafa Keles, MD</td>
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<td>“Living Donation Education”</td>
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<tr>
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<tr>
<td>Social Media / Facebook Apps</td>
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**References**

Papers of particular interest, published recently, have been highlighted as:

- Of importance
- Of major importance


Figure 1.
Used with permission from the Clinical Journal of American Society of Nephrology.
Figure 2.
“House Calls” or Home-Based Education Program
Talking About Live Kidney (TALK) Donation and TALK-Social Worker Program

Video and booklet on LDKT and living donation
Those materials plus sessions with a social worker

Measure:
probability of patient doing any of the following

> discuss living donation with family or physicians

> initiate or complete transplant evaluation

> identify a potential living donor

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<th>Usual Education</th>
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<td>30%</td>
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Figure 3.
Talking About Live Kidney (TALK) Donation and TALK-Social Worker Program
Living ACTS

Videos and booklets about LDKT and living donation were presented to African American patients in a culturally sensitive way.

Figure 4.
Living ACTS
Culturally Targeted Website for Hispanic Transplant Candidates

Patients and their families had access to a bilingual, culturally targeted website about LDKT and living organ donation

Knowledge gains for patients with access to the website were substantially more than for those without access

Figure 5.
Culturally Targeted Website for Hispanic Transplant Candidates
Seminars on Saturdays

On Saturdays, talk with and provide material to patients and family and friends, targeted toward preemptive kidney transplantation

> Preemptive LDKT from non-blood relatives increased from 25% to 58%

> Percentage of non-blood related donors among total living donors increased from 28% to 41%

Figure 6.
Seminars on Saturdays
Live Donor Champion

Transplant candidates identify a family member or friend to be trained as an advocate. Advocates share information about the benefits of LDKT and the patient's need for an organ with their social network.

My friend needs a kidney donor

Pilot study found increases in living donor evaluations and LKDT

**Figure 7.**
Live Donor Champion
Figure 8.
Social Media/Facebook Apps

Social Media/Facebook Apps

Facebook smartphone app enables transplant candidates to share their need for a living donor through social media.

Patients using the app were nearly 7 times as likely to have a potential donor come forward.
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<th>Program</th>
<th>Additional Program Description</th>
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| “House Calls” or Home-Based Education Program, (Rodrique et al, 2007 & 2014) | • HB intervention in which trained health educators perform 60 to 90-minute LDKT education session with family and members of the candidate’s social network.  
• Sessions may be held in an alternative setting (e.g., another residence, local library) if the patient is uncomfortable hosting the session or if a larger setting is needed to accommodate more guests.  
• Goals are to (a) increase transplantation and living donation knowledge and awareness among members of the patient’s social network who are unable or unlikely to attend clinic-based educational sessions, and (b) jump-start the discussion about LDKT and living donation.  
• For most members of the patient’s social network, HB education may be their first opportunity to learn about LDKT, and to have their questions answered by a transplant health educator. More knowledge among social network members may facilitate discussions about LDKT and reduce patients’ hesitancy to discuss the topic with others.  
• HB education for black patients included at least 1 minority educator and incorporation of race-specific transplant statistics into discussion, and written brochures highlighted minority recipients and their donors.  
• In the first (~1 hour) session, the candidate’s social network is diagrammed in order to facilitate understanding of the candidate’s social system, and to guide the candidate in formulating a list of invitees to the second session.  
• In the second (~2.5 hour) session, which includes the transplant candidate’s social network, the educators provide information on kidney disease, dialysis, LDKT, and living donation; stimulate open communication between the patient and family members; and seek to develop consensus on the patient’s goals and how the goals could be achieved with engagement and support of the social network. |
| “Kidney Team at Home” (Ismail et al, 2014; Massey et al, 2016) | • In this adaptation of “House Calls”, the HB intervention comprises 2 visits at the transplant candidate’s home. The first visit (1 hour) focuses solely on the patient, while the second visit (2 to 3 hours) includes the transplant candidates’ social network and explores the possibilities of LDKT. |
| “Talking About Live Kidney Donation” (TALK) and “TALK-Social Worker” Intervention Programs (Boulware et al, 2013; DePasquale et al, 2012) | • The 20-minute video includes ethnically diverse patients and families describing their experiences discussing and pursuing LDKT, and transplant professionals describing factors for patients/families to consider.  
• The booklet encourages patients not yet on dialysis to talk to family and health care providers about LDKT, with examples of model conversations and sources of additional information. Patients may review the video and booklet on their own or with a social worker.  
• Patients are encouraged to share the materials with members of their social network.  
• In the first “TALK-Social Worker” session, the social worker distributes the video and booklet and helps the patient identify barriers to considering or pursuing LDKT. The patient is asked to invite family/friends to a second session.  
• At the second session, the social worker helps social network members to identify barriers to LDKT and living donation and to describe prior discussions about living donation.  
• Problem-solving and motivational interviewing techniques are employed at both sessions. |
| “Living ACTS” (About Choices in Transplantation and Sharing) (Arriola et al, 2014) | • The program was based on a conceptual model that emphasizes the need to address both “surface” characteristics relevant to a given cultural group (e.g., by including people, places and language familiar to and preferred by the target audience) and more deeply rooted elements that may influence behavior (e.g., the influential roles that family discussion and family impact play in many African-Americans’ decisions regarding healthcare).  
• The video features African-American patients, families and healthcare professionals and addresses the impact of LDKT on families, how family decision making around living donation may occur, and financial resources for LDKT.  
• The booklet provides additional information such as web links and tips for starting conversations with family members. The intervention was designed to address inadequacies of existing educational efforts in addressing unique concerns and issues in African American patients that may serve as barriers to pursuit of LDKT. |
**Program** | **Additional Program Description**
---|---
"Infórmate" Culturally Targeted Website (Gordon et al, 2015 & 2016) 16,17 |  
- Focus groups with Hispanic kidney transplant candidates, living donors, dialysis patients, and members of the general public supported content development.
- This culturally-targeted website is presented in Spanish and English and was designed to extend the Northwestern Memorial Hospital's Hispanic Kidney Transplant Program, which is a unique clinical program providing care to Hispanic transplant patients.

"Written Educational Materials followed by Structured Educational Session" (Barnieh et al, 2011) 18 |  
- This session, which includes 3 to 5 other patients and their invited family members, uses a problem-based learning format and is facilitated by a transplant nurse coordinator and nephrologist.
- In addition to providing didactic information about the advantages and disadvantages of ESRD treatment options and living kidney donation, patients and family members form smaller groups to problem-solve common scenarios pertaining to LDKT. For instance, this includes finding strategies to overcome barriers to living donation and role playing how to ask others about living donation.
- At the end of the session, there is a question and answer session with a LDKT recipient and former living kidney donor.

ESRD, end-stage renal disease; HB, home-based; LDKT, living donor kidney transplantation
**Table 2**
Description of programs supported by observational (non-randomized) studies

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| “Patient Navigators in Community-Based Nephrology Practices” (Marlow et al, 2016) | • The patient navigator intervention is predicated on the assumption that early contact with transplant candidates allows for early education about transplantation and a higher potential for LDKT, particularly preemptive transplants.  
• The overarching goal of the program is to help patients progress through the referral, evaluation, and transplantation process.  
• Social workers seek to establish relationships with potential transplant candidates, provide support and individualized transplant and living donation information to patients and family members, and address barriers to possible LDKT and living donation. |
| “Living Donor Family Education” (Schweitzer et al, 1997 & 2000; Foster et al, 2002) | • In addition to a didactic presentation and discussion led by one of several program nurse coordinators, an 8 to 10-minute video incorporates information on the surgical procedure and risks, as well as perspectives and comment from donors on the pre-operative testing experience.  
• The video seeks to give perspectives from donors demographically similar to the population served by the transplant center (predominantly African American and older).  
• Overall, the program seeks to address problems in the transplant center’s past educational efforts, including: 1) limited and inconsistent provision of information on living donation to transplant candidate families; 2) lack of integration of education into the program’s evaluation and care process. |
| “Seminars on Saturdays” (Brown et al, 2011) | • This program represents a revision of an existing program offered by a single transplant center, based on attendee feedback and analysis of logistical barriers felt to have limited previous participation in education programs.  
• The program is conducted by a multidisciplinary team including surgeons, other physicians, and nursing staff including transplant coordinators, clinical nurses, pre-dialysis nurse specialists, dialysis staff, counselors, and 10–12 kidney recipients and donors (per seminar).  
• The inclusion of testimonials from laypersons, who varied in age/gender/ethnicity and type relationship to the patient, is viewed as a critical component of the program.  
• The program is offered every 2 months (6 times per year) on a Saturday to all pre-dialysis patients and their family/friends, as well as to all patients on the kidney waitlist, and to all dialysis patients not yet waitlisted but deemed suitable for transplant by their care providers.  
• The program seeks to address the following problems: 1) lack of access to information and misconceptions about living donation, both of which identified as principal barriers to pursuit of LDKT by patients; 2) a need to improve the quality and delivery of information. |
| “Live Donor Champion” (LDC) (Garonzik-Wang et al, 2012) | • The LDC program focuses on removing three specific barriers to LDKT: 1) discomfort discussing such a sensitive subject; 2) fear of asking someone else for an organ, and 3) limited knowledge about the live donation process.  
• The role of the LDC is to improve comfort in initiating conversations and spreading awareness about the patient’s kidney failure and about live donation.  
• Anyone over age 18 who speaks English is eligible to serve as a LDC; the LDC is usually a spouse, significant other, family member, or friend of the patient.  
• The program comprises 6 meetings held once monthly. Each 2-hour meeting focuses on a different topic. Topics include introduction to kidney transplantation and living donation, how to initiate a conversation with potential living donor candidates, identifying a social network, and sharing success stories with a living donor and recipient panel.  
• The LDC is encouraged to use the information and skills learned at each session to begin looking for potential donors. Participants are asked to keep a monthly log of their progress and are also encouraged to record what worked well and what did not to provide feedback to program coordinators. |
| “Hispanic Transplant Education” (Gordon et al, 2014) | • The program sought to address some of the underlying reasons for disparities in LDKT rates in Hispanics.  
• The first session includes both patients and family members (or friends) in which a Hispanic transplant surgeon presents information about transplantation, donation risks and benefits, and alternatives. |
<table>
<thead>
<tr>
<th>Program</th>
<th>Additional Program Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>“Educational Intervention Based in Peer Experiences” (Delair et al, 2010)²⁶</td>
<td>• The second session includes only family members and is focused on cultural concerns and misconceptions.</td>
</tr>
<tr>
<td></td>
<td>• Exposure to the experience of living liver donors is not usually included in transplant candidate education; the investigators hypothesized that it could help transplant patients to improve their knowledge about living donation and to feel more comfortable discussing donation with others.</td>
</tr>
<tr>
<td></td>
<td>• In the materials, the former living liver donors discuss their experiences, including views about the surgery, recovery, costs, employment, and life after donation. The materials also include direct quotes from donors concerning what information they would give to individuals considering living liver donation.</td>
</tr>
<tr>
<td></td>
<td>• Although the intervention is designed to be delivered to the transplant candidate in the transplant clinic, it can be mailed to patients.</td>
</tr>
<tr>
<td></td>
<td>• Patients are encouraged to share the materials with potential living donors in their social network.</td>
</tr>
<tr>
<td>“Pre-Dialysis Education” (PDEP) (Cankaya et al, 2013)²⁷</td>
<td>• An implicit hypothesis was that PDEP would have an impact on preemptive LDKT.</td>
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<tr>
<td></td>
<td>• The modules address general considerations and treatment recommendations for all patients regardless of level of kidney disease; and peritoneal dialysis, hemodialysis and kidney transplantation.</td>
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<tr>
<td></td>
<td>• The problems that the PDEP was designed to address included the need to 1) inform and educate patients about their disease and treatment options before dialysis, and 2) reduce treatment complications, increase treatment compliance, and start treatment in a planned way to avoid emergencies.</td>
</tr>
<tr>
<td>“Living Donation Education” (LED) (Cervera et al, 2015)²⁸</td>
<td>• The intervention is delivered in two phases at the transplant center.</td>
</tr>
<tr>
<td></td>
<td>• In Phase I, patients and family members attend a formal education session in which a healthcare provider progresses through an audiovisual presentation on alternative ESRD treatments, the benefits of LDKT, living and deceased donation, specific information focused on living donation (eligibility criteria, surgery, risks, outcomes), the concept of a ‘live donor champion’, and strategies to identify potential living donors.</td>
</tr>
<tr>
<td></td>
<td>• All participants receive a booklet mirroring information presented during the session.</td>
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<td></td>
<td>• In Phase II, patients and family members are invited to meet with a living donor and recipient pair who share their experiences and answer questions.</td>
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<tr>
<td></td>
<td>• Both educational sessions and the booklet are provided in English or Spanish, based on patient preference.</td>
</tr>
<tr>
<td>Social Media/ Facebook Apps (Kumar et al, 2016)²⁹</td>
<td>• Grounded in prior literature, clinical judgement, ethicist oversight and feedback from patient focus groups, the smartphone app provides step-by-step instructions for creating a Facebook post detailing an individual’s struggle with organ failure and their need for a living donor.</td>
</tr>
<tr>
<td></td>
<td>• Links to sources of vetted information on the risks, benefits and processes of live donation are also included.</td>
</tr>
<tr>
<td>“My Transplant Coach” Decision Aid (Axelrod et al, 2017)³⁰</td>
<td>• Developed by educational professionals, medical professionals, statisticians and patient representatives with feedback from focus groups, the app provides patient-centric information about kidney transplantation to improve overall transplant knowledge and allow candidates to consider the relative benefits of LDKT and various types of DDKT options.</td>
</tr>
<tr>
<td></td>
<td>• Incorporates data based on waitlist mortality, geographic variation in organ supply and survival calculators.</td>
</tr>
<tr>
<td></td>
<td>• Presents data to candidates using animated presentation and simple graphics.</td>
</tr>
</tbody>
</table>

LDC, live donor champion; LDKT, living donor kidney transplantation
### Table 3
Common Design Elements of Effective Living Donor Transplant Education Programs

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><strong>Include a patient’s family and friends.</strong> When family and friends learn with the patient, more conversations happen about the patient’s need, transplant options, and living donation.</td>
</tr>
<tr>
<td>2</td>
<td><strong>Conduct in a comfortable place.</strong> Patients may learn more readily in places other than a doctor’s office such as library, a community center or a home.</td>
</tr>
<tr>
<td>3</td>
<td><strong>Provide written materials or other media.</strong> Providing patients with simply stated factual material about the benefits and risks of living donation and living donor transplantation increases their understanding and comfort.</td>
</tr>
</tbody>
</table>
### Table 4
Core Content of Living Donor Transplant Education Programs

<table>
<thead>
<tr>
<th>Benefits of transplantation</th>
<th>Living donor evaluation process</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average waiting times for transplantation</td>
<td>Living donor eligibility criteria</td>
</tr>
<tr>
<td>Transplant evaluation process</td>
<td>Living donor surgery</td>
</tr>
<tr>
<td>Types of transplants and donors</td>
<td>Typical donor recovery</td>
</tr>
<tr>
<td>Transplant outcomes</td>
<td>Risks of living donation</td>
</tr>
<tr>
<td>Current deceased donation rates</td>
<td>Benefits of living donation</td>
</tr>
<tr>
<td>Differences in donation rates (living and deceased) by race (for minority patients)</td>
<td>Common donor concerns</td>
</tr>
<tr>
<td></td>
<td>Common recipient concerns</td>
</tr>
<tr>
<td></td>
<td>Possible indirect costs of donation process</td>
</tr>
<tr>
<td></td>
<td>Helpful resources for transplant candidates</td>
</tr>
<tr>
<td></td>
<td>Helpful resources for living donors</td>
</tr>
</tbody>
</table>
### Table 5

Common Delivery Teams for Living Donor Transplant Education Programs

<table>
<thead>
<tr>
<th></th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><strong>Transplant Professionals.</strong> Educational program delivery teams include diverse professionals such as surgeons, social workers, doctors, nurses, and others.</td>
</tr>
<tr>
<td>2</td>
<td><strong>Living Donors.</strong> Involvement of living donors creates opportunities for patients and members of their support network to talk with living donors about their decision, surgery, recovery and subsequent quality of life.</td>
</tr>
<tr>
<td>3</td>
<td><strong>Transplant Recipients.</strong> Opportunities for patients to talk with prior transplant recipients who have been through the process of identifying a living donor reduces hesitancy and empowers patients to talk about living organ donation.</td>
</tr>
</tbody>
</table>