December 22, 2017

Patrick Mauro
Contract Specialist
Department of Health and Human Services (HHS)
Health Resources and Services Administration (HRSA)
5600 Fishers Lane
Rockville, MD 20857
Submitted via email: pmauro@hrsa.gov

Re: Organ Procurement and Transplantation Network
Solicitation Number: 18-250-SOL-00017

Dear Mr. Mauro:

On behalf of the American Society of Transplant Surgeons (ASTS), we are pleased to have this opportunity to comment on the Organ Procurement and Transplantation Network (OPTN) Performance Work Statement (the “RFI”). ASTS is a medical specialty society representing approximately 1,800 professionals dedicated to excellence in transplantation surgery. Our mission is to advance the art and science of transplant surgery through leadership, advocacy, education, and training.

Our comments primarily focus on those provisions of the RFI that relate to the OPTN’s role in the oversight of transplant centers and data collection.

**OPTN Oversight of Transplant Centers**

The RFI includes a number of Tasks related to review and periodic reassessment of membership applications (3.6.1), ongoing review of Transplant Centers through the Membership and Professional Standards Committee (MPSC) (3.2.5.6), and monitoring of member performance (3.6). Historically, the OPTN has engaged in substantial regulatory oversight over Transplant Centers, through the development of detailed policy, survey processes, MPSC reviews, and data reporting requirements. Much of this activity is duplicative of regulatory review conducted by the Centers for Medicare and Medicaid Services (CMS) through implementation of the Transplant Center Conditions of Participation (CoPs).
**Recommendation:** We request that the RFI be modified to include a new Task requiring the Contractor to engage in a joint effort with HRSA and CMS to eliminate the duplicative regulatory oversight of Transplant Centers.

The objective of this effort should be to institute a regulatory oversight process that includes:

- One set of outcomes criteria
- One set of regulations and interpretive guidelines
- One jointly conducted survey, and
- One set of consequences for those centers that fail to meet applicable requirements.

Our more detailed proposal setting forth options for eliminating unnecessary duplication in Transplant Center regulation and oversight is attached as **Appendix A**. Since the involvement of the OPTN will be necessary to make meaningful progress on these objectives, we request that the RFI include an additional Task that requires Contractor involvement in this effort.

We note that NOTA does not delegate to the OPTN any responsibility over organ transplantation. The functions of the OPTN, as specifically listed in NOTA, include the requirement that the OPTN “adopt and use standards of quality for the acquisition and transportation of donated organs,” but no mention is made of OPTN authority to regulate the transplantation of those organs.

Moreover, the NOTA regulations related to the OPTN (42 CFR § 121.1 et. seq.) (the Final Regulation) does not anticipate duplicative OPTN regulation of Transplant Centers that are approved for participation in Medicare and that are in compliance with the CMS CoPs. Specifically, the Final Regulation (42 CFR §121.3(b)) states, in relevant part:

(b) **Membership of the OPTN.**

(1) The OPTN shall admit and retain as members the following:

(ii) Transplant hospitals participating in the Medicare or Medicaid programs;

(Emphasis added.) The information required to be submitted by a transplant hospital in order to be admitted as a member of the OPTN is specified by regulation and limited to the name and address of the hospital and a list of its transplant programs by organ.1 While the OPTN does have the authority to deny the membership applications of entities other than OPOs and transplant hospitals, it does not appear that the OPTN has the authority to deny membership to either OPOs or transplant hospitals that apply.2

---

1 42 CFR §121.3(b)(2)).
2 A transplant hospital may be excluded from membership and a designated transplant program may be precluded from receiving organs if there is a risk to the health of patients or public safety, under the procedures set forth at 42 CFR §121.10(c).
Section 121.9(a) of the Final Regulation elaborates on the requirements that must be met for a transplant program to be eligible to receive organs for transplantation (a “designated transplant program”). This provision sets forth three different routes for a transplant program to be considered a “designated transplant program”: (1) The program may “be a transplant program approved by the Secretary for reimbursement under Medicare”; OR (2) The Program is one which has “adequate resources to provide transplant services to its patients” and meet certain requirements as specified by the OPTN related to the scope and breadth of services that must be provided by the transplant center; OR (3) The program may be in a Department of Veterans Affairs (VA), Department of Defense (DOD), or other Federal hospital. Thus, under the plain regulation of the Final Regulation, only those transplant programs that are not participating in the Medicare program and are not VA, DOD or other Federal hospitals are required to meet requirements set forth in OPTN policies (such as, for example, policies pertaining to operating and recovery room resources, intensive care resources and surgical beds and transplant program personnel; policies requiring evidence of collaborative involvement of experts in various fields of medicine (e.g. infectious disease, radiology), requirements related to access to certain laboratory services and the availability of psychiatric and social support services). The Final Regulation clearly suggests that transplant centers that are approved for Medicare participation are not required to meet OPTN requirements with regard to these and similar operational issues.

Moreover, the Final Regulation provision that addresses the OPTN’s policy-making role does not anticipate the OPTN’s imposing additional performance requirements on Medicare-participating Transplant Centers. The only specific transplant-center related policies that the Final Regulation directs the OPTN Board to establish are: (1) policies regarding the training and experience of transplant surgeons and physicians in designated transplant programs (42 CFR §141.4(a)(4)) and (2) procedures for transplant hospitals to make reasonable efforts to obtain financial resources for patients unable to pay for transplantation and transplant follow-up (42 CFR §141.4(a)(3)(ii). Current OPTN policies imposing various operational requirements on Medicare certified Transplant Centers significantly exceed those specifically required to be adopted under the Final Regulation.

Finally, the provision of the Final Regulation related to reviews, evaluation, and enforcement does not anticipate duplicative oversight of Medicare participating Transplant Centers, except to the extent that such a Transplant Center’s performance raises a risk to the health of patients or public safety, in which case the HHS may require the OPTN to take action under the procedures set forth at 42 CFR §121.10(c). We believe that much of the OPTN oversight of Transplant Centers is a historical holdover from the period before adoption of the Medicare CoPs. Before CMS adopted the CoPs in 2007, the OPTN was necessarily much more involved in determining the quality and other standards that Transplant Centers should meet. Prior to the adoption of the CoPs, CMS only required Transplant Centers to be an OPTN member and to perform a minimum number of transplants meeting specified outcomes requirements in order to participate in the Medicare program. When CMS adopted CoPs in 2007, however, substantial redundancy and inconsistency was introduced into the regulatory system. Now, over ten years later, the time has come to eliminate this duplicative regulatory structure. Since the OPTN’s involvement is integral to any effort to reduce or eliminate duplicative regulation of Transplant Centers by the OPTN/HRSA and CMS, we believe that the RFI should include this as a separate Task in the OPTN Contractor’s scope of work.
**Data Collection**

The RFI includes a number of requirements related to data collection. Significantly, for example, Requirement 3.5 requires the OPTN Contractor to consider all data collected by the Contractor to be “official OPTN data” that must be collected through OMB approved data collection forms, unless an exception applies under the Paperwork Reduction Act. The “Performance Standard” for this task requires the Contractor to submit OMB data collection forms for all OPTN data not currently collected through OMB process.

As noted in our comments on prior OPTN RFIs, the OPTN currently imposes extensive data requirements on Transplant Centers, and it is possible that OMB data collection forms have not been submitted for some or all of these data elements. We are also concerned that requiring OMB data collection forms for all data elements reduces the OPTN’s ability to collect needed data without unnecessary administrative delay and could prevent timely collection of important data.

**Recommendation:** *We request that this Task be modified to require the OPTN to work with its members to determine which of these data elements impose an unnecessary administrative burden and which are needed for outcome evaluation, risk adjustment, or other critical OPTN functions.*

**General Observations**

ASTS would like to take the opportunity presented by the RFI to urge HRSA to use the RFI to refocus the OPTN’s activities more narrowly on its basic mission, as set forth in NOTA and the Final Regulation. As originally enacted, NOTA specifically lists the functions to be carried out by the OPTN:

(A) establish in one location or through regional centers "(i) a national list of individuals who need organs, and "(ii) a national system, through the use of computers and in accordance with established medical criteria, to match organs and individuals included in the list, especially individuals whose immune system makes it difficult for them to receive organs,

(B) maintain a twenty-four-hour telephone service to facilitate matching organs with individuals included in the list,

(C) assist organ procurement organizations in the distribution of organs which cannot be placed within the service areas of the organizations,

(D) adopt and use standards of quality for the acquisition and transportation of donated organs,

(E) prepare and distribute, on a regionalized basis, samples of blood sera from individuals who are included on the list and whose immune system makes it difficult for them to receive organs, in order to facilitate matching the compatibility of such individuals with organ donors,
(F) coordinate, as appropriate, the transportation of organs from organ procurement organizations to transplant centers,

(G) provide information to physicians and other health professionals regarding organ donation, and

(H) collect, analyze, and publish data concerning organ donation and transplants.

Over the years, the mission of the OPTN has expanded into virtually every aspect of transplantation and the operations of the OPTN have been encumbered by bureaucratic processes, obfuscating the importance of its basic mission. For example, the Performance Work Statement—which is basically an outline of the OPTN’s tasks—is 68 pages long, and includes the following tasks for the Contractor to perform:

- 3 general requirements,
- 10 contract administration requirements,
- 19 requirements pertaining to governance and committees,
- 3 requirements pertaining to policy development,
- 8 requirements pertaining to data collection,
- 11 requirements pertaining to access to OPTN data,
- 6 requirements pertaining to communication with the SRTR,
- 4 requirements pertaining to communications,
- 1 requirement related to reports to Congress,
- numerous requirements related to data security and privacy,
- 1 requirement related to records management training for OPTN employees,
- 1 requirement pertaining to transition-in planning,
- 1 requirement to transition-out planning, and
- 1 requirement related to special studies (content unspecified).

By contrast, the RFI lists only six requirements directly pertaining to the OPTN’s matching function.

**Recommendation:** We respectfully suggest that HRSA streamline and minimize the paperwork requirements imposed on the OPTN, and substitute tasks that focus more closely on substantive tasks that have the potential to further improve the expeditious and equitable distribution of organs.

The mission elements of the OPTN seem to us to be out of order: increasing the number of transplantable organs takes a back seat to micromanaging transplant center operations.
**Recommendation:** We respectfully suggest that HRSA refocus the OPTN on saving lives through organ transplantation and donation and place “Working actively to increase the supply and utilization of donated organs” first in the list of mission elements. We also suggest that HRSA include creating a plan to increase organ donation in the OPTN tasks.

The current contractor has worked with the transplant community to develop a strategic plan which we believe should be reflected in the Performance Work Statement.

Finally, we would like to note that the distribution of the RFI and timeline for response appeared more limited than in the past. We hope that you will not perceive a limited response as lack of interest in the RFI and the role the community can play in enhancing the work of the OPTN. We appreciate the opportunity to submit these comments on the RFI. If you have any questions about these comments, please do not hesitate to contact ASTS Executive Director Kim Gifford via phone (703-414-1609) or email (kim.gifford@asts.org).

Sincerely yours,

Jean C. Emond, MD
President
Via e-mail

September 11, 2017

Thomas E. Price, MD
Secretary of Health and Human Services
Hubert H. Humphrey Building
200 Independence Avenue S.W.
Washington, D.C. 20201

Re: American Society of Transplant Surgeons Request for Regulatory Relief

Dear Secretary Price:

On behalf of the American Society of Transplant Surgeons, we applaud your initiative to minimize regulatory burdens on physicians that interfere with the efficient and effective delivery of high quality care. Along these lines, we strongly believe that this initiative should address the current duplicative regulation of transplant centers (TCs) by both the Centers for Medicare and Medicaid Services (CMS) and the Organ Procurement and Transplantation Network (OPTN) under the auspices of the Health Resources and Services Administration (HRSA). This letter sets forth an overview of the problem, a proposed framework for addressing it, and proposed next steps.

I. Overview

A. In General

For the past several years, the transplant community has been facing a steadily mounting burden of oversight that is now threatening the creative and vibrant spirit that has marked the field since its inception, like the frog in the old adage sitting in slowly heating water until it finds itself boiling. We recognize the complexity of our enterprise and embrace the challenge of caring for our patients with compassion and expertise. We also welcome the opportunity to seek ever-improving results in a safe and reliable patient care system. We are proud to publicly demonstrate our outcomes and our ever-improving processes to ensure patient safety and fairness to all who need a transplant. However, despite the culture of pride and innovation that permeates our community, individual infractions by a small number of transplant programs have led to an overwhelming burden of oversight on transplantation and transplantation-related care.
Under current law, both the OPTN and CMS impose both process and outcomes requirements on TCs. The CMS and OPTN outcomes requirements differ, and for that reason the TCs identified for review by the OPTN and those identified as out of compliance with Medicare approval requirements differ. In addition, the OPTN and CMS impose different process requirements on transplant centers, and their methods of ensuring compliance with several of the common requirements differ. A crosswalk of CMS and OPTN requirements available on the OPTN website suggests that, together, there are approximately 123 requirements (a number that we believe to be underestimated), approximately 30% of which are reviewed by both CMS and the OPTN. The remaining requirements relate principally to Quality Assurance & Performance Improvement (QAPI) and multi-disciplinary team requirements imposed by CMS but not the OPTN, differing review processes for pediatric and adult programs, differing volume (clinical experience) requirements, and differing waitlist management and notification requirements. Both the CMS and the OPTN regulatory processes require extensive review of medical records. This duplication of regulatory requirements is costly both for TCs and for the federal government (and its paid private contractors), and unnecessarily distracts time and attention from patient care. Preparation for a CMS or OPTN survey is a months-long process requiring hundreds of FTE hours as well as considerable physical resources. Both surveys require an egregious waste of paper, when it could all likely be done electronically. Moreover, both sets of requirements are overly prescriptive and interfere with the patient-physician relationship and with physician judgment in the context of complex clinical decision-making.

The imposition of these extraordinarily burdensome regulatory requirements is particularly inappropriate in light of the extremely demanding outcomes requirements imposed by both the OPTN and CMS. Both CMS and the OPTN impose strict patient and graft one-year survival requirements that dissuade TCs from accepting “suboptimal” organs. Post-transplant organ graft and patient survival expectations are set significantly higher for transplantation than other diseases like cancer, which has a 66.9% five-year survival rate. As the chart below illustrates, transplantation truly has excellent outcomes and ideally should be made available to a greater number of recipients.¹

<table>
<thead>
<tr>
<th>Organ</th>
<th>One-Year</th>
<th>Three-Year</th>
<th>Five-Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart</td>
<td>91%</td>
<td>89%</td>
<td>85%</td>
</tr>
<tr>
<td>Intestine</td>
<td>81</td>
<td>80</td>
<td>67</td>
</tr>
<tr>
<td>Kidney</td>
<td>97</td>
<td>96</td>
<td>93</td>
</tr>
<tr>
<td>Liver</td>
<td>91</td>
<td>89</td>
<td>83</td>
</tr>
<tr>
<td>Lung (Single and Double)</td>
<td>87</td>
<td>85</td>
<td>69</td>
</tr>
<tr>
<td>Pancreas</td>
<td>92</td>
<td>93</td>
<td>88</td>
</tr>
<tr>
<td>Heart-Lung</td>
<td>80</td>
<td>73</td>
<td>59</td>
</tr>
<tr>
<td>Kidney-Pancreas</td>
<td>98</td>
<td>96</td>
<td>95</td>
</tr>
</tbody>
</table>

In light of the excellent outcomes achieved by TCs, it is unclear to us why such extraordinarily onerous “process” requirements are considered necessary by not only one government agency, but two.

**B. Some Examples**

The overlapping (and sometimes conflicting) morass of TC regulatory requirements imposed by both CMS and the OPTN are ripe for simplification and streamlining. Not only are these requirements duplicative, they are also extraordinarily detailed. For example:

- The OPTN regulatory requirements include 230 pages of policy, 180 pages of Bylaws, and 65 pages of Evaluation Plan.  
  
  2 https://optn.transplant.hrsa.gov/media/1200/optn_policies.pdf  

- CMS regulatory requirements includes 85 pages of the Federal Register, 107 pages of the Survey and Certification Interpretive Guidelines, 50 pages of Survey and Certification Interpretive Guideline Changes, 49 pages of Quality Assessment and Performance Improvement (QAPI) Program requirements, 103 pages of updated Interpretive Guidelines (pending revisions May 2016), and numerous additional updates and clarifications.  
  
  3 See, e.g.  

- **Organ and Vessel Tracking**
  - The time out and verification process has been cumbersome. Requirements in this area have been characterized by variation in surveyor preference between the OPTN and CMS and by each agency’s practice of changing its own regulatory requirements, in an asynchronous manner over time. Forms developed by the OPTN do not meet the
requirements of CMS surveyors, and CMS and OPTN surveyors have imposed different requirements regarding electronic documentation in the Electronic Medical Record (EMR) versus paper copy and signatures.

- **Patient Education and Consent to Proceed**
  - It is critical that patients and living donors are well informed of the risks, benefits, and alternatives to participation in the various phases of transplant and donation. However, regulations call these encounters “Informed Consent” which create unnecessary concern from the patient and confusion with legal and risk departments within a hospital. The OPTN consent requirements are incredibly prescriptive yet they do not provide a templated form for use. TCs find themselves constantly updating the forms to include small language changes, only to then be cited for missing something minuscule. Meeting these informed consent requirements has resulted in extremely lengthy documents and encounters to “cover all bases,” which are overwhelming to both recipients and potential donors, and which add significant physician time.

- **Time Requirements and Disparate Timeframes**
  - The OPTN and CMS are on separate timelines for completing onsite and offsite surveys. Often TCs encounter both teams within a given year, resulting in 3-5 days of interruption for each visit. OPTN surveys are scheduled and CMS surveys are unscheduled. These agencies do not coordinate TC survey schedules, which has resulted in disruption of TC hospital and clinical operations. Survey readiness, even when scheduled, also necessitates significant TC financial and resource commitment. In addition, significant team time is required to implement plans of correction, education, and auditing after the visit. Surveys are useful tools when they result in process improvements; however, typically these surveys result in more administrative work, redundant documentation, and team time away from clinical care. In one large center, over 60 staff are occupied full time at the expense of patient care and other duties the week of the CMS survey.

- **Clinical Micromanagement**
  - Process oversight extends to the micromanagement of educational materials provided to patients and specific documentation requirements of the role of social workers and others in the overall care of patients in multiple locations. For example, the transplant nephrologist must document the results of the psychiatric assessment even when the consultation report of the psychiatrist is part of the patient’s chart.

- **ABO Verification**
  - A single highly publicized misallocation of a heart by blood type led to a process of verification of blood types prior to transplantation that has grown into a morass of checklists and forms, which are dated and timed and avidly reviewed by surveyors for spelling errors and marks in the wrong box. Up to 63 points of failure have been identified in form completions and design. Compliance with this regulation has led to a substantial number of new administrative positions to oversee the work of the clinicians and evaluate form completion and requirements. While well intentioned, these processes do not contribute to patient safety, only paperwork and administrative burden.
C. Statutory Authority

Despite the extraordinary level of detailed oversight imposed by both CMS and the OPTN, neither Medicare certification nor OPTN TC review processes are clearly or unequivocally required by statute. Section 1881(b)(1) of the Social Security Act (the “Act”) gives the Secretary authority to prescribe regulations for payment for renal transplantation services; however, there does not appear to be any specific statutory authority mandating certification of renal transplant programs, and no provision of the Act of which we are aware mandates the establishment of any type of specialized certification requirements for other forms of transplantation. In fact, CMS only adopted specific certification regulations for TCs in March of 2007, as the result of earlier public and Congressional concerns raised in response to certain highly publicized lapses by a handful of TCs. Before that time, CMS relied entirely on the OPTN to oversee TCs, since Section 1138 (a)(1)(B) of the Social Security Act, enacted in 1986, requires Medicare and Medicaid participating hospitals that perform organ transplants to be members of and abide by the rules and requirements of the OPTN. CMS relied on its general rulemaking authority to publish rules and regulations “necessary for the efficient administration of the functions” of the Medicare Program to adopt the final TC certification regulations, which reflects the lack of specific statutory authority to establish TC certification requirements independent of OPTN membership criteria.

The OPTN was established by the National Organ Transplant Act (NOTA) (PUBLIC LAW 98-507-OCT. 19, 1984), and Section 372(b)(2) of NOTA sets forth with specificity the responsibilities of the OPTN. None of these explicitly requires the OPTN to establish quality requirements for TCs; however, NOTA does indicate that the OPTN has the responsibility to establish its own membership standards, and it appears that this is the sole statutory basis for the OPTN’s extensive oversight of TCs. While it may be argued that comprehensive oversight of TCs by the OPTN was necessary when the OPTN was the only body charged with ensuring that TCs maintained quality standards, since the adoption of Medicare certification standards, that is no longer the case.

II. A Proposed Framework to Reduce TC Regulatory Burden

A. Principles to Guide TC Regulatory Reform

Because TC regulatory requirements are imposed by two separate and independent agencies within HHS and because the requirements imposed by both agencies are detailed and complex, administrative simplification in this arena may prove challenging. For this reason, the administrative simplification process should be guided by clear and easily understood basic principles, the objective of which is to preserve transplant quality and patient protections, while simplifying and streamlining oversight. Specifically, we strongly believe that the regulatory review process should result in:

- One set of TC oversight regulations and regulatory interpretation;
- One set of TC outcomes measures intended to maximize transplantation rates;
- One combined survey conducted as necessary based on a single set of survey triggers; and
- One set of consequences for noncompliance.

We refer to these objectives as the “Reform Principles.”

4 42 USC §274(b)(2)(B)
B. Operationalizing the Reform Principles

The Reform Principles could be operationalized in any number of ways. As discussed above, currently the OPTN and CMS oversee and monitor both outcomes and process. However, it may be possible to apply the Reform Principles by providing one of the two agencies with the authority to establish outcomes requirements while the other establishes process requirements, or, following historical areas of special competence, the OPTN might be given primary responsibility for establishing the requirements for activities and processes that occur outside the four walls of the TC (e.g., organ retrieval, allocation and distribution, patient ranking on the waitlist, and ensuring the fairness of waitlist processes), while CMS retains primary authority to establish rules related to TC activities, including QAPI (an area in which CMS and its contractor have established special experience and expertise).

However, in our experience, each of the two agencies has considerable expertise that the other does not, and the ideal regulatory framework would involve close collaboration of CMS and the OPTN/HRSA to establish a single integrated regulatory framework and oversight process. For this reason, we urge the Secretary to consider a TC regulatory framework with the following characteristics:

- **Approval of New TCs:** Currently, CMS does not regulate TCs until they become operational, and the job of approving new centers falls to the OPTN. We believe that this allocation of responsibility is appropriate and that, while the OPTN requirements for new centers should be reviewed and streamlined to the extent practicable, the OPTN should retain the responsibility for initial TC approval.

- **Organ Retrieval and Allocation, Waitlist Management, and Related Data Management:** Likewise, the OPTN has considerable expertise in the area of waitlist management and oversight, and has comprehensive processes in place to ensure that waitlist rules are not subject to “gaming.” In addition, the OPTN routinely engages in considerable data collection to ensure compliance with organ allocation and distribution policies. In operationalizing the Reform Principles, we urge the Secretary to direct the OPTN/HRSA to review its current standards related to these and other areas that take place outside the “four walls” of the TC, but to retain OPTN/HRSA sole oversight authority in these areas. To the extent that on-site surveys must be conducted to ensure compliance with such waitlist, allocation or other rules, the survey should be conducted as part of a unified OPTN/HRSA/CMS survey (discussed below).

- **Establishment of Interagency Committee:** We urge the Secretary to appoint an interagency committee (“Interagency Committee”) composed of representatives appointed by CMS, HRSA, and the OPTN, to operationalize the Reform Principles and to enforce compliance. The tasks of the Interagency Committee should include at least the following:
  - **Unified Outcomes Requirements:** Under current rules, TCs’ one-year outcomes, as reported and risk adjusted by the Scientific Registry of Transplant Recipients (a HRSA contractor), are assessed by CMS and by the OPTN using different statistical standards, resulting in the “flagging” of different centers and different times by the two agencies. We urge the Secretary to direct CMS and the OPTN/HRSA to establish a single set of TC outcomes standards and to modify Medicare certification regulations and/or OPTN Bylaws and policies as necessary to adopt the agreed outcomes standards.
Streamlined Process Requirements Focused on Transparency and Due Process: TCs are among the only, if not the only, type of provider, other than Organ Procurement Organizations (OPOs) that are required to comply with both outcomes requirements and comprehensive process requirements as a condition of participation in the Medicare Program. Each TC publicly reports its individual patient and organ survival statistics, and, under current Medicare certification standards, a TC that reports lower than expected patient and graft survival for two years fails to meet Medicare certification standards. We do not believe that TCs that maintain high outcomes standards also should be subject to comprehensive process requirements related to various aspects of clinical care. Since outcomes standards are generally viewed as out-of-reach for many types of providers, Medicare certification requirements generally focus on compliance with processes thought to contribute to positive outcomes. In the absence of outcomes standards that are definable and enforceable, the imposition of process requirements is viewed as the “best that can be done.” Where, as in the case of TCs, outcomes standards are available and applied on a continual basis, why should CMS or OPTN also impose comprehensive and detailed process requirements related to various aspects of clinical care? On the other hand, there are certain transparency and due process standards that all TCs should maintain regardless of the outcomes they achieve. We believe that it is in these areas that process requirements and oversight surveys should focus:

- Patient rights
- Core safety measures
- QAPI programs
- Care of living donors

Non-Prescriptive Interpretive Guidelines: Currently, much of the burden of compliance with both CMS and OPTN regulatory requirements arises not because of the basic CMS regulations and OPTN Bylaws or policies, but because of the overly prescriptive manner in which these requirements are applied bysurveyors from the two agencies. The Interpretive Guidelines used by surveyors under contract with CMS are currently over 100 pages in length, and an equally long revision is currently on hold as the result of substantial objections from the TC community. Likewise OPTN surveyors utilize a comparable 65 page document, the OPTN Evaluation Plan, in conducting OPTN surveys, of approximately equal length. We urge the Secretary to direct the Interagency Committee to develop a single interpretive document to be used by those conducting surveys of compliance with those relatively limited process requirements that are retained, and that, in developing this document, the Interagency Committee should be requested to use the following guidelines:

- The Interagency Committee should develop Interpretive Guidelines with the clear objective of minimizing administrative burden. Care should be taken to ensure that the Interpretive Guidelines stay within the scope of limited process requirements.

---

The Interpretive Guidelines should be developed with input from the transplant community. When CMS and the OPTN both have current interpretive documents relating to the same basic requirement, the least prescriptive guideline should be adopted as the model. To the extent that TCs are expected to utilize a particular form or process, a model document should be provided to the transplant community.

- Triggers for Oversight Survey: Oversight surveys should be performed in a coordinated fashion and should involve surveyors from both CMS and the OPTN, working together. Surveys should be performed under well-defined circumstances, such as:
  - Failure to meet outcomes thresholds
  - Major sentinel events
  - Failure to comply with OPTN rules regarding listing and allocation practices

- Consequences of Failure to Comply with Process/Meet Outcomes Standards: The Interagency Committee should merge the OPTN and CMS plan of correction/mitigating circumstances processes; review of TC performance should be conducted jointly by the OPTN/SRTR and CMS; and the process should draw upon the expertise of the OPTN Membership and Professional Standards Committee (MPSC).

III. Next Steps

The reforms outlined above likely would require modification of the CMS Conditions of Participation for TC and OPTN Policies (and possibly Bylaws). It is also possible that the task of streamlining the TC review process should be included in the scope of services set forth in the Request for Proposals to be issued by HRSA for the OPTN contract, which we understand will be released this Fall.

We would be delighted to meet with CMS, OPTN, and HRSA representatives to discuss the next steps in developing a unified, streamlined, and effective TC oversight process. If we can provide any further information regarding our concerns or proposed framework for addressing the overregulation of TCs, please do not hesitate to contact Kim Gifford, ASTS Executive Director, at kim.gifford@asts.org or 703-414-7870.

Sincerely yours,

Jean C. Emond, MD
President
American Society of Transplant Surgeons