Review

Sodium Iodide Symporter for Nuclear Molecular Imaging and Gene Therapy: From Bedside to Bench and Back

Byeong-Cheol Ahn

Department of Nuclear Medicine, Kyungpook National University School of Medicine and Hospital, Daegu, South Korea

Corresponding author: Byeong-Cheol Ahn, MD, PhD, Professor and Director, Department of Nuclear Medicine, Kyungpook National University School of Medicine and Hospital, 50 Samduck 2-Ga, Jung Gu, Daegu, South Korea 700-412. Tel: 82-53-420-5583; Fax: 82-53-422-0864; E-mail: abc2000@knu.ac.kr

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Received: 2011.10.28; Accepted: 2012.01.23; Published: 2012.04.11

Abstract

Molecular imaging, defined as the visual representation, characterization and quantification of biological processes at the cellular and subcellular levels within intact living organisms, can be obtained by various imaging technologies, including nuclear imaging methods. Imaging of normal thyroid tissue and differentiated thyroid cancer, and treatment of thyroid cancer with radioiodine rely on the expression of the sodium iodide symporter (NIS) in these cells. NIS is an intrinsic membrane protein with 13 transmembrane domains and it takes up iodide into the cytosol from the extracellular fluid. By transferring NIS function to various cells via gene transfer, the cells can be visualized with gamma or positron emitting radioisotopes such as Tc-99m, I-123, I-131, I-124 and F-18 tetrafluoroborate, which are accumulated by NIS. They can also be treated with beta- or alpha-emitting radionuclides, such as I-131, Re-186, Re-188 and At-211, which are also accumulated by NIS. This article demonstrates the diagnostic and therapeutic applications of NIS as a radionuclide-based reporter gene for trafficking cells and a therapeutic gene for treating cancers.

Key words: sodium iodide symporter, molecular imaging, radionuclide-based imaging, gene therapy, radionuclide.

Introduction

The ability of the thyroid gland to concentrate iodide has long provided the basis for diagnosis and therapeutic management of benign thyroid diseases and thyroid cancer [1]. Thyroid scintigraphy with radioiodines or technetium-99m (Tc-99m) pertechnetate has played a key role in the evaluation of thyroid nodules with its ability of providing anatomical and functional information since the advent of modern endocrinology [2]. Radioiodine via an 'atomic cocktail' was first used medically for thyroid cancer treatment under the Atomic Energy Act since 1946 [3]. Thereafter, millions of patients with benign or malignant thyroid diseases have been given radioiodine for diagnostic and therapeutic purposes with successful outcomes. However, the uptake mechanism of radioiodine into thyroid tissue or thyroid cancers was not fully elucidated until 1996, when the sodium iodide symporter (NIS) was finally cloned [4]. This not only improved understanding of thyroid pathophysiology tremendously, but also offered promising molecular biological strategies of imaging and treatment. Clinical theranostic application of NIS function using radioiodine was projected to biologic preclinical experimental studies after the NIS cloning. NIS expression can be imaged feasibly with simple radiotracers, such as radioiodines or Tc-99m. By the easy imagina-
ble characteristic of NIS, it has been used as an imaging reporter to monitor gene transfer.[5, 6] In addition to the potential as the imaging reporter gene, NIS has been used as a therapeutic gene to treat cancers through its ability to concentrate therapeutic doses of radionuclides in target cells [7-10].

This review is mainly focused on the theranostic application of NIS for radionuclide-based molecular imaging and radionuclide gene therapy in in vivo animal models.

NIS

NIS is an intrinsic plasma membrane glycoprotein with 13 transmembrane domains that actively mediates iodide transport into the thyroid follicular cells and several extrathyroidal tissues [11]. This protein plays an essential role in thyroid physiology by mediating iodide uptake into the thyroid follicular cells, a key step in thyroid hormone synthesis. NIS belongs to the sodium/solute symporter family or solute carrier family 5, which drives negatively-charged solutes into the cytoplasm using an electrochemical Na⁺ gradient [12]. The symporter co-transporters two sodium ions (Na⁺) along with one iodide (I⁻), with the transmembrane sodium gradient serving as the driving force for iodide uptake; therefore, NIS functionality is dependent on the electrochemical sodium gradient that is maintained by the ouabaine-sensitive Na⁺/K⁺-ATPase pump (Fig. 1) [13].

NIS needs to be localized in the plasma membrane for efficient transportation of iodide into thyroid follicular cells. Poor iodide uptake in thyroid cancer cells compared to thyroid follicular cells is related to impaired targeting and retention of NIS at the membrane. Membrane localization of NIS requires thyroid stimulating hormone (TSH) stimulation; through TSH deprivation, NIS is not retained at the membrane, leading to a decrease in iodide uptake. Although TSH stimulation is essential for efficient NIS trafficking to plasma membrane of thyroid follicular cells, it is possible that TSH-independent mechanisms for the trafficking exist because non-thyroidal tissues also retain NIS at the membrane in the absence of TSH stimulation. One suggested mechanism of NIS targeting to the membrane is the phosphorylation of NIS at serine residues in the carboxy terminus. Protein-protein interaction is another suggested mechanism for the trafficking. NIS contains PDZ, dileucine and dipeptide motifs which might be associated with trafficking [1, 13]. Non-thyroidal cancer tissues also can express NIS; however, only 20-25% of NIS-positive tumors showed iodide uptake partly due to the intracytoplasmic location of NIS [14].

Although expression of NIS is also detectable in normal extrathyroidal tissues such as the salivary glands, gastric mucosa and lactating mammary glands, the expression is not regulated by TSH and is present at lower levels in these tissues than in thyroid tissue. Iodide organification is a particular and unique characteristic of the thyroid gland, and long-term retention of iodide does not occur in the extrathyroidal tissues expressing NIS [15].

![Figure 1](http://www.thno.org/attachment.png) Figure 1. Iodide uptake function of NIS. NIS transports 2 sodium ions and 1 iodide ion into the cytoplasm together. The electrochemical sodium gradient generated by the ouabaine-sensitive Na⁺/K⁺-ATPase pump provides energy for this transfer.

Radiopharmaceuticals for NIS

NIS has marked advantages as an imaging reporter gene and as a therapeutic gene compared to other reporter or therapeutic genes due to the wide availability of radiopharmaceuticals and its well understood metabolism and clearance of these radiopharmaceuticals from the body [16].

NIS actively takes up radioiodide and Tc-99m; therefore, its function can be imaged with I-123, I-131, I-124 and Tc-99m [7, 15, 17]. No issues of labeling processes and stability arise when using these radiopharmaceuticals, whereas they may be a major concern of the radiolabeled ligands of other radionu-
side-based reporter genes, such as the dopamine D2 receptor or herpes simplex virus thymidine kinase (HSV-tk) genes [16].

I-123 is produced in a cyclotron by proton irradiation of enriched xenon-124 (Xe-124) in a capsule, decays by electron capture to tellurium-123 (Te-123) with a half-life of 13.2 hours, and emits gamma rays with predominant energies of 159 keV (the gamma ray is primarily used for imaging) and 127 keV. I-123, mainly a gamma emitter, has a high counting rate compared with I-131 and provides a higher lesion-to-background signal, thereby improving sensitivity and imaging quality. Moreover, with the same administered activity, I-123 delivers an absorbed radiation dose that is approximately one-fifth that of I-131 to NIS-expressing tissues [18].

I-124 is a proton-rich isotope of iodine produced in a cyclotron by numerous nuclear reactions and decays to Te-124 with a half-life of 4.2 days. Its modes of decay are 74.4% electron capture and 25.6% positron emission. It emits gamma radiation with energies of 511 and 602 keV [19].

I-131 is produced in a nuclear reactor by neutron bombardment of natural Te-127, decays by beta emission with a half-life of 8.0 days to Xe-133, and emits gamma rays as well. It most often (89% of the time) expends its 971 keV of decay energy by transforming into the stable Xe-131 in two steps, with gamma decay following rapidly after beta decay. The primary emissions of I-131 decay are beta particles with a maximal energy of 606 keV (89% abundance, others, 248–807 keV) and 364 keV gamma rays (81% abundance, others 723 keV) [19]. As I-131 emits both beta and gamma rays, it can be used to image NIS gene expression; however, it is not recommended for imaging due to poor image quality (by high energy of the gamma rays) and the high radiation burden (by the beta rays) compared to I-123.

Tc-99m, a metastable nuclear isomer of Tc-99, has a half-life of 6.0 hours and emits 140 keV gamma rays which is an optimal energy for scintigraphic imaging. Tc-99m, the most commonly used radionuclides in routine nuclear medicine imaging, is usually extracted from Tc-99m generators which contain parent nuclide molybdenum-99 (Mo-99) [2].

Recently, F-18 tetrafluoroborate (F-18 TFB) was developed as a positron-emitting radiopharmaceutical that is actively taken up by NIS [20]. The rapid uptake and efflux of F-18 TFB in the rat thyroid cell line parallels the behavior of Tc-99m pertechnetate, which is known to be taken up in cells expressing NIS [20]. Uptake of F-18 TFB to thyroid follicular cells is stimulated by TSH and blocked by perchlorate. It was suggested that F-18 TFB transport occurs with little or no coupling to sodium transport, or that TFB occupies a binding site on NIS but is transported very inefficiently.

I-131, rhenium-188 (Re-188), Re-186 and astatine-211 (At-211), which emit particles from their nuclei, are used for radionuclide therapy on cells expressing NIS [1, 8-10, 13, 21]. Re-188 is an important therapeutic radionuclide, which is obtained on demand as a carrier-free sodium perrhenate by saline elution of the tungsten-188 (W-188)/Re-188 generator system. With a half-life of 17.0 hours and emission of a high-energy beta ray (maximal energy of 2.12 MeV) and gamma ray (155 keV, 15%) for imaging, Re-188 offers the prospect of cost-effective preparation of radiopharmaceuticals for cancer treatment [22]. Cyclotron-driven neutron activator may be an alternative for on-demand supply of Re-188 [23].

Currently, At-211 is the most promising alpha-emitter that has been studied for cancer therapy. It is the heaviest halogen, with no stable isotope. It decays via a double-branch pathway with a mean alpha-energy of 6.7 MeV (42% 5.9 MeV and 58% 7.5 MeV) and a half-life of 7.2 hours. As a consequence of its electron capture branching to its daughter polonium-211, X-rays of 77 to 92 keV in sufficient abundance are emitted, enabling external imaging (including single photon emission computed tomography [SPECT]) and gamma counting of blood samples as additional advantages. However, its widespread use in therapeutic doses is hindered as a result of limited availability of medium-energy cyclotrons with an alpha-particle beam for its production, which is currently feasible at only a few research centers [24]. Table 1 summarizes characteristics of radionuclides which can be used with NIS for diagnostic or therapeutic purposes.

Use of NIS for diagnostic purposes in clinical nuclear medicine

Gamma camera imaging with radioiodine (I-131 or I-123) can visualize metastatic lesions in differentiated thyroid cancer patients who have undergone total thyroidectomy because the lesions are highly efficient at trapping circulating iodine by expression of NIS (Fig. 2) [25]. Radioiodine scintigraphy, once the mainstay of post-therapy imaging surveillance, has largely been replaced by neck ultrasonography as the modality of choice for long-term imaging surveillance, although it still may be used for the detection of occult or distant metastases, particularly in the setting of a newly elevated serum thyroglobulin level [26]. Routine use of radioiodine scintigraphy for surveillance is not recommended for low-risk patients. However, it is still used in patients with intermediate...
or high risk of recurrence, as well as to assess patients for evidence of recurrence in the setting of an elevated thyroglobulin level with a negative neck ultrasonography. Scintigraphy performed after empiric treatment with high doses of I-131 is more sensitive than the usual diagnostic I-131 scanning [26].

I-124 positron emission tomography (PET) has higher sensitivity for the detection of thyroid cancer lesions with NIS expression compared with I-131 whole body scintigraphy due to lower background noise and the higher resolution of PET imaging than gamma camera imaging. Additionally, PET images can be fused with CT and/or magnetic resonance imaging [27].

Detection and localization of metastatic thyroid cancer lesions by radiiodine scintigraphy or PET rely on the expression of NIS in the cancer cells which accumulate radioiodine [27].

### Table 1. Radionuclides used for diagnostic or therapeutic purposes associated with NIS.

| Radioisotopes | T_{1/2}   | Emission          | Most abundant energy | Medical use                        |
|--------------|-----------|-------------------|-----------------------|-----------------------------------|
| I-123        | 13.2 hours| gamma ray         | 159 keV               | Diagnostic imaging (Gamma camera) |
| I-124        | 4.2 days  | positron/gamma ray| 511 keV/602 keV       | Diagnostic imaging (PET)          |
| I-131        | 8.0 days  | beta ray/gamma ray| 606 keV/364 keV       | Diagnostic imaging (Gamma camera) Therapy |
| Tc-99m       | 6.0 hours | gamma ray         | 140 keV               | Diagnostic imaging (Gamma camera) |
| F-18         | 109.8 minutes | positron         | 511 keV               | Diagnostic imaging (PET) as a form of tetra-fluoroborate |
| Re-186       | 90.6 hours| beta ray/gamma ray| 1,070 keV/59 keV      | Therapy                           |
| Re-188       | 17.0 hours| beta ray/gamma ray| 2,120 keV/155 keV     | Therapy                           |
| At-211       | 7.2 hours | alpha ray/X-ray   | 7,500 keV/77-92 keV   | Therapy                           |

T_{1/2}: half-life, PET: positron emission tomography

**Figure 2.** A 21-year-old female who underwent total thyroidectomy due to papillary thyroid cancer. Chest simple radiography and CT did not demonstrate any metastatic lesion of the cancer in the neck and chest regions. However, a radiiodine whole body scan revealed lymph node metastases (white arrow) in the right supraclavicular area and diffuse lung metastases (black arrows).
NIS and nuclear molecular imaging

Molecular imaging, defined as the visual representation, characterization and quantification of biological processes at the cellular and subcellular levels within intact living organisms, can be obtained by various imaging technologies, such as optical imaging, nuclear imaging, magnetic resonance imaging (MRI), ultrasound imaging and computed tomography (CT) [28]. Molecular imaging has the potential to provide unique information that will guarantee the safety and efficacy of biotherapies which utilize antibodies, bacteria or cells in humans, and also will contribute to the future development of novel biotherapies [15].

With the emergence of cell therapies in regenerative medicine, it is important to track cells injected into subjects. In this context, NIS has been used in preclinical studies. With transfer of the NIS gene into therapeutic cells such as cytotoxic T or natural killer cells, nuclear molecular imaging modalities can image the cells with a relevant radiotracer, such as I-123, I-124, I-131, Tc-99m or F-18 TFB. The NIS-expressing cells have been imaged with planar scintigraphy, SPECT or PET according to the administered radio-tracers [15].

Nuclear imaging modalities, such as PET and SPECT, provide the 3-dimensional distribution of radiopharmaceuticals and have excellent sensitivity and high resolution with excellent tissue penetration depth [29]. These advantages permit these imaging techniques for use in translational research, from cell culture to preclinical animal models to clinical applications [28]. Both PET and SPECT give quantitative and non-invasive information on NIS gene expression or the number of NIS-expressing cells [15, 28].

As a gene reporter, NIS is able to be used for monitoring of gene and vector biodistribution and for trafficking of therapeutic cells [6, 15]. Contrary to the diagnostic application of radioiodine nuclear imaging using NIS gene expression for the detection of thyroid cancer recurrence or metastases, NIS gene transfer is a prerequisite for radionuclide-based molecular imaging (Fig. 3). Non-invasive imaging of NIS expressing nonthyroidal cells with a gamma camera or PET upon viral gene transfer has been demonstrated feasible and safe in experimental animals and humans as well (Fig. 4) [6, 8].

Figure 3. Cells without NIS gene expression obtain the function of iodine uptake with NIS gene transduction by viral or non-viral vector delivery. The cells can be imaged by radionuclide-based molecular imaging techniques using gamma ray or positron-emitting radiotracers and be cleared by beta or alpha particle-emitting radionuclides.
Figure 4. Visualization of macrophages expressing NIS with radionuclide-based molecular imaging. Inflammation at the right thigh (yellow arrow) was well visualized in F-18 FDG microPET imaging. Migration of macrophages expressing NIS to the inflammation site (white arrow) was clearly visualized on I-124 microPET imaging [7].

Figure 5. Visualization of tumor cells expressing NIS with optical molecular imaging using I-124. Tumor xenografts of anaplastic thyroid cancer cells expressing NIS were well visualized on both microPET imaging (white arrows) and Cerenkov luminescence imaging (black arrows) after intravenous administration of I-124 [17].

Recently, I-131 and I-124, which are commonly used for thyroid imaging, were reported to have sufficient energy to result in Cerenkov radiation that can be visualized with sensitive optical imaging equipment and cells transfected with NIS gene were successfully imaged with the radiiodines using an optical imaging instrument in an in vivo animal model (Fig. 5) [17].

Radioiodine accumulation in NIS-expressing organs such as the thyroid is a deterrent to scintigraphic visualization of NIS-expressing cells in various animal models. To remove radioiodine uptake in the thyroid gland and better visualize NIS-expressing cells, the animal can be prepared with surgical total thyroidectomy or radioiodine ablation before administration of the NIS-expressing cells [30].
Use of NIS for therapeutic purposes in clinical nuclear medicine

Molecular radionuclide-based therapy of differentiated NIS-expressing thyroid cancer with I-131 was the cornerstone on which nuclear medicine was built and it has been a very successful example of targeted therapy to reduce recurrence and mortality for almost 70 years (Fig. 6) [31, 32]. Therapeutic application of I-131 for hyperthyroidism and thyroid cancer was implemented in the early 1940s, and success of the applications resulted in the approval of medical radioisotope use and initiation of atomic medicine, later re-named nuclear medicine [31, 33].

Radioiodine therapy for thyroid diseases relies on the fact that thyroid follicular cells and differentiated thyroid cancer are efficient at trapping circulating radioiodine than other tissues [25]. I-131 treatment has been the most preferred therapeutic modality by physicians for hyperthyroidism in the United States and it has been one of the key treatment modalities for differentiated thyroid cancers worldwide [34, 35]. However, I-131 treatment is not very effective in de-differentiated thyroid cancer, which down-regulates NIS expression, and is meaningless in anaplastic thyroid and medullary thyroid cancers, which do not express NIS. One possible treatment option for de-differentiated thyroid cancer is the induction of re-differentiation with differentiating agents such as retinoic acid and thiazolidinedione [31, 36].

Expression of NIS is not uncommon in breast and stomach cancers, and some reports have shown visualization of primary or metastatic lesions of such cancers with radiiodine or Tc-99m scintigraphy [37-40]. The possibility of radiiodine treatment for cancers with sufficient NIS expression has been suggested; however, as far as the author knows, clinical reports on such treatment with successful outcome have yet to be published, likely due to insufficient NIS expression [16].

Although it has not been clinically attempted, anaplastic or medullary thyroid cancers lacking NIS expression can be treated with I-131 after NIS gene transfer to the tumors. Additionally, other tumor entities which do not express NIS can also be treated with the same strategy [31].

Figure 6. A 26-year-old female who underwent total thyroidectomy due to papillary thyroid cancer. (A) Chest simple radiograph did not demonstrate any observable metastatic lesions of the cancer. (B) CT scan of the chest demonstrated several metastatic lesions of the cancer in both lung fields (white arrows). TSH-stimulated serum thyroglobulin was 65.0 ng/mL. The patient was diagnosed with metastatic thyroid cancer of the lung. (C) A post initial high dose I-131 treatment (150 mCi) scan revealed numerous metastatic lung lesions. (D) A post 2nd high dose I-131 treatment (200 mCi) scan revealed fewer but still several metastatic lung lesions (black arrows). (E, F) A post 3rd high dose I-131 treatment (200 mCi) scan revealed no remarkable radiiodine uptake in both lung fields and chest CT showed only tiny lung nodules having no clinical significance. TSH-stimulated serum thyroglobulin was 1.4 ng/mL after the third treatment. The patient had achieved complete remission with three times of high dose I-131 treatment and her status still remains disease-free at 7 years follow-up.
NIS and radionuclide gene therapy

In addition to its imaging potential, NIS can be used as a therapeutic gene through its ability to concentrate therapeutic doses of radionuclides in target cells [15]. Contrary to the therapeutic application of I-131 using NIS gene expression for treating thyroid cancer recurrence or metastases, NIS gene transfer is a prerequisite for tumors without NIS gene expression. After the transfer of the NIS gene into various cancer cells, they can be treated with beta or alpha particle-emitting radionuclides including I-131, Re-186, Re-188 and At-211, which are accumulated via NIS (Fig. 3) [31].

Right after NIS was cloned by Carrasco et al. in 1996, many researchers started to use the gene for therapeutic purposes with I-131, and in general, the results were effective. The effect of I-131 NIS gene therapy was enhanced with higher doses of I-131 and intervention with retinoic acid or dexamethasone, which increase radioiodine uptake [41]. Transcription factors such as Pax-8 and TTF-1 could induce or promote iodide uptake and specifically prolong iodide retention time in cancer cells [42, 43].

Re-188 and At-211 were also used as therapeutic radionuclides with NIS gene therapy to nonthyroidal tumors. Re-188 has advantages over I-131, as its beta ray energy is higher and has a shorter half-life, which makes it a more suitable radionuclide for NIS-expressing tumors. In addition, it is conveniently obtained from a W-188/Re-188 generator [44]. At-211, which emits extremely cytotoxic alpha-particles, is known to be taken up by NIS in thyroid tissue and has been used as a therapeutic radionuclide for NIS-expressing tumors in cell culture and animal experiments [21, 45, 46]. In addition to very effective tumoricidal effects, At-211 has the advantages of alpha-particle’s short range and a short half-life, which allow for a minimal radiation burden to the surrounding environment, including people [46].

However, single radionuclide NIS gene therapy might have limited therapeutic effects and can produce serious adverse effects positively related to the amount of administered radionuclide dose. Reducing the radionuclide dose for NIS gene therapy is able to reduce the adverse effects, but might lead to limited effectiveness [47]. Combined treatment of radionuclide NIS gene therapy with other therapeutic approaches could be more efficient to improve therapeutic outcomes and can reduce adverse effects of radionuclide NIS gene therapy by reducing the radionuclide dose. Chemotherapy, genciclovir HSV-tk gene therapy, immunotherapy, external beam radiotherapy and siRNA therapy have been combined with radionuclide NIS gene therapy, and the results were almost always successful [8-10, 47].

Even though radionuclide NIS gene therapy has been shown to be effective in in vivo animal models, several issues must be resolved before this novel strategy can be useful clinically. First of all, vector systems having safe, effective and specific NIS gene delivery to the tumor are needed. The optimal time interval between NIS gene transfer and therapeutic radionuclide administration should be determined to obtain the most effective therapeutic results. Organs that normally express NIS, such as the thyroid gland and the salivary glands, are inevitably damaged by the therapeutic radionuclide; therefore, protecting or managing strategies for the organs need to be developed [13, 31, 48].

Even though radionuclide NIS gene therapy is only performed in the preclinical setting at the moment, clinical trials of the treatment are likely to happen in the not-too-distant future with advances in efficiency and safety of the therapy by close communication between these basic biological studies and clinical experiences of thyroid cancer treatment with I-131.

NIS-based molecular imaging and radionuclide gene therapy; limitations and future directions

NIS provides an advantage of both as reporter and therapeutic genes and therefore, NIS gene transfer makes it possible to image, monitor and treat the tumor with appropriate radionuclides, just as in differentiated thyroid cancer. Another advantage of NIS is wide availability of appropriate diagnostic and therapeutic radiopharmaceuticals. Although NIS is one of the best theranostic genes, there are several pending questions that must be answered before its clinical use.

Tissues that normally express endogenous NIS such as the thyroid gland, salivary glands and stomach, are an obstacle for NIS-based imaging or treatment. Uptake of imaging radiotracers to the tissues conceals trafficking target cells expressing NIS which are located near the tissues. Uptake of therapeutic radionuclides to the normal tissues can damage the organs and may reduce tracer uptake to the target cells expressing exogenous NIS.

Retention time of radioiodine is generally short in NIS-transduced cells by rapid washout of the radiiodine, therefore absorbed dose and toxicity to the target cells might be limited and it precludes successful radioiodine NIS gene therapy. To prolong the retention time, drugs such as lithium carbonate, or...
co-transfer of the thyroid peroxidase gene was introduced; however, results were conflicting and not very effective [13]. Co-transfer of the thyroglobulin gene was also suggested to increase retention time [42]. Efflux of iodine from the cell is known to be related to pendrin, SLC5A8 and ClCn5, and even though not verified by experiments, down-regulation of these proteins can delay iodine efflux from the cell [42]. Ablation of the thyroid gland and low iodine diet are able to prolong the retention time in NIS transduced tumor cells, however applicability of this strategy is limited in a clinical situation. It can be feasibly applied only in thyroid cancer patients receiving previous thyroidectomy. Enhancement of radiiodine uptake by up-regulation of NIS expression has been tried with drugs such as retinoic acid or dexamethasone, troglitazone and external radiation [49, 50]. Histone deacetylase inhibitors (e.g. depsipeptide, trichostatin A and valproic acid) and demethylating agents (5-azacytidine) have been used to restore endogenous NIS expression [1]. In addition to increasing radiation dose to the NIS expressing cells, radiosensitization can enhance the biological effect of the same radiation dose. DNA damage repair inhibitors revealed a therapeutic benefit with radionuclide NIS gene therapy [51]. Further studies are needed for validation and optimization of the pharmacological approaches for prolonging the retention time, delaying iodine efflux, restoring/up-regulation of NIS expression and enhancing radiosensitization before practical use.

Several new diagnostic or therapeutic radiopharmaceuticals for NIS were recently studied. Cells expressing NIS can be imaged with F-18 FTB PET instead of radiiodine scintigraphy and be treated more effectively with Re-188, Re-186 or At-211 instead of I-131. Some of the radiopharmaceuticals are not suitable at present due to scarce availability and nontrivial safety issues related to their production and handling. Technical advancement of the production and handling skills for the radiopharmaceuticals is warranted.

With administration of I-131, the thyroid gland takes up I-131 and retains it within the gland for a long time by organification of the radiiodine. This will end in permanent hypothyroidism by radioablation of normal thyroid tissue. The salivary gland also accumulates the radionuclide and xerostomia can occur by radiation sialoadenitis related to uptake of the radionuclide. To maintain sufficient radiiodine uptake to the extrathyroidal cancer tissues expressing NIS, uptake of radiiodine to the thyroid gland can be suppressed by thyroid hormone replacement and antithyroidal drugs [52]. Stable iodine administration before administration of radiiodine can reduce radiiodine to the gland as well [13]. Radiiodine uptake in the salivary gland can be expelled by manual massage of the gland and may reduce incidence of xerostomia related to radiation-induced sialoadenitis [53]. Strategies for preventing or reducing side effects to normal tissues expressing NIS by radionuclides uptake must be developed and optimized before common clinical application of NIS-based radionuclide theranostics.

Conclusions

Although diagnostic and therapeutic use of the NIS gene began in clinics more than half a century ago, understanding of the biology of NIS has been advancing rapidly in the last two decades. NIS-based molecular imaging and radionuclide gene therapy, cutting edge technologies in molecular imaging and gene therapy arenas, were born with imitation of diagnostic and therapeutic applications in the field of clinical thyroid practice. With fast advancement of molecular imaging and gene therapy with active research, these bench technologies are likely to be used in the clinical setting in the near future.

Acknowledgements

This study was supported by a grant (A102132) of the Korea Health Technology R&D Project, Ministry of Health & Welfare, Republic of Korea and the Ministry of Knowledge Economy (MKE), and a grant of the Korea Institute for Advancement of Technology (KIAT) and Daegyeong Leading Industry Office through the Leading Industry Development for Economic Region.

Competing Interests

The authors have declared that no competing interest exists.

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