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Prostate Cancer Evaluation: Design, Synthesis and Evaluation of Novel Enzyme-Activated Proton MRI Contrast Agents

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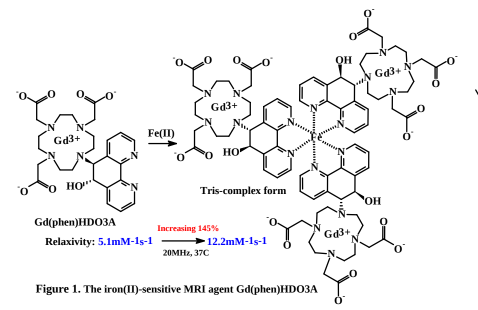
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### **INTRODUCTION**

**BACKGROUND** Prostate cancer is the most frequently diagnosed cancer and the second leading cause of cancer death in men in the United States, in 2009, approximately 192,000 men were diagnosed with prostate cancer with 27,000 succumbing to this disease,[1,2] currently, there is no cure for locally advanced or metastatic prostate cancer.

Gene therapy has emerged as a potentially promising strategy for treatment of prostate cancer.[3-15] The prostate is particularly amenable to gene therapy.[11-16] However, there are major issues in terms of assessing the delivery to target tissue, assessing the uniformity (versus heterogeneity) of biodistribution and determining whether the genes are expressed.[15-33] A viral construct is often readministered on successive occasions, but this should optimally be timed to coincide with loss of expression. Inevitably gene therapy has associated risks, and thus non-invasive in vivo determining the duration of gene expression in an individual tumor could greatly enhance the viability of the approach. Gene expression now is commonly monitored by in situ hybridization techniques or by introducing a marker gene to follow the regulation of a gene of interest. Since  $\beta$ -galactosidase ( $\beta$ -gal) activity is readily assessed by histology or in culture, in hosts as evolutionarily diverse as bacteria, yeast, and mammals, its introduction has become a standard means of assaying clonal insertion, transcriptional activation, protein expression, and protein interaction, *lacZ* gene encoding *E. coli* β-gal has already been recognized as the most commonly used reporter system.[34] A variety of lacZ gene reporters has been developed, such as colorimetric, [35-39] fluorescence, [40-53] chemiluminescence, [54-61] radiotracers for positron emission tomography (PET) or single-photon emission computed tomography (SPECT),[62-66] magnetic resonance imaging (MRI) probes, [67-69] and <sup>19</sup>F-NMR approaches, [70-77] though most of them have only been utilized in in vitro detection, with a very few successful applications in vivo so far.[39,49,50,51,60,63-65,67,68,76,77] Therefore, the development of non-invasive lacZ gene reporter techniques based on appropriate molecules and imaging modalities is still a high desire.

The superb spatial resolution and the outstanding capacity of differentiating soft tissues have determined the widespread success of magnetic resonance imaging (MRI) in clinical diagnosis.[78] The contrast in an MR image is the result of a complex interplay of numerous factors, including the relative  $T_1$  and  $T_2$  relaxation times, proton density of the imaged tissues and instrumental parameters. It was shown that contrast agent causes a dramatic variation of the water proton relaxation rates, thus providing physiological information beyond the impressive anatomical resolution commonly obtained in the uncontrasted images. Contrast agents are widely used clinically to assess organ perfusion, disruption of the blood-brain barrier, occurrence of abnormalities in kidney clearance, and circulation issues.[78-82] The responsive MRI contrast agents holds great promise in the gene therapy arena.[83,84] The abilities of these contrast agents to relax water protons is triggered or enhanced greatly by recognition of a particular biomolecule opening up the possibility of developing MRI tests specific for biomarkers indicative of particular disease states and aiding in the early detection and diagnosis of tumors. Desreux et al [80,85] demonstrated that, by chelating Gd(phen)HDO3A with Fe(II) to form a highly stable triscomplex, as shown in Figure 1, the relaxivity increased 145% at 20MHz and 37°C from 5.1mM<sup>-1</sup>s<sup>-1</sup> per Gd(III) in Gd(phen)HDO3A form to 12.2 mM<sup>-1</sup>s<sup>-1</sup> in the tris-complex. Desreux *et al* [80,85] also synthesized another iron-sensitive MRI contrast agent with a tris-hydroxamate (Figure 2). After the trishydroxamate groups formed a chelate with Fe(III), free rotation at the Gd(III) centers was restricted, thereby increasing relaxivity by 57% from 5.4 to 8.5mM<sup>-1</sup>s<sup>-1</sup> at 20 MHz.



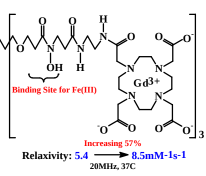
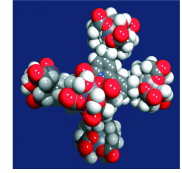


Figure 2. The iron(III)-sensitive MRI agent Gd(III)-Trishydroxamic acid

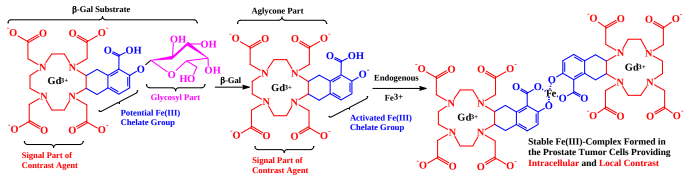
Most recently, Merbach *et al* [86-88] observed the remarkably high  $T_1$  relaxivity gain by the heterometallic, self-assembled metallostar formation with six efficiently relaxing GdIII centers from (tpy-DTTA)Gd(H<sub>2</sub>O) with 7.3mM<sup>-1</sup>s<sup>-1</sup> to {Fe<sup>II</sup>[Gd<sup>III</sup><sub>2</sub>(tpy-DTTA)<sub>2</sub>(H<sub>2</sub>O)<sub>4</sub>]<sub>3</sub>}<sup>4-</sup> with 15.7mM<sup>-1</sup>s<sup>-1</sup> at 20MHz and 37°C (**Figures 3**), significantly, their detailed studies on structure and dynamics of the trinuclear complex {Fe<sup>II</sup>[Gd<sup>III</sup><sub>2</sub>(tpy-DTTA)<sub>2</sub>(H<sub>2</sub>O)<sub>4</sub>]<sub>3</sub>}<sup>4-</sup> indicate that the heterometallic self-assemblies attain high  $T_1$  relaxivities by influencing three factors: water exchange, rotation, and electron relaxation.



**Figure 3.**  $\{Fe[Gd_2L_2(H_2O)_4]_3\}^{4-}$ 

**DESIGN** Prompted by these findings, we proposed a novel class of enzyme responsive Gd<sup>3+</sup>-based MRI contrast agent for high sensitivity and specificity for  $\beta$ -gal detection, based on the tumor biology and Fe-chelation therapeutic strategy. Cancer cells, as compared with their normal counterparts, exhibit increased uptake and utilization of more iron, as evidenced by an increase in transferrin receptors at the cancer cell surface, mediating a high level and rate of iron uptake.[89] More recently, an emerging class of Fe-chelator agents have shown effective antitumor activity *in vitro* and *in vivo*, which can overcome resistance to standard chemotherapy, due to their ability to affect multiple molecular targets including the enzyme responsible for the rate-limiting step of DNA synthesis, ribonucleotide reductase, molecules involved in cell cycle control (*e.g.* cyclin D1, p21<sup>CIP1/WAF1</sup>) and the inhibition of metastasis (*i.e.* N-myc downstream regulated gene-1).[89-94] The FDA has approved five Fe-chelators for use in anticancer therapy so far, some others are in clinical trials for the treatment of various metastatic and solid cancers.[89,95-97] In our design, the *lacZ* responsive Gd<sup>3+</sup>-based MRI contrast agent is comprised of three moieties: (A) a signal enhancement group, such as Gd-DOTA or Gd-PCTA; (B) an Fe<sup>3+</sup> chelating group; (C)  $\beta$ -D-galactose. Upon encountering with  $\beta$ -gal in tumor cells, the released, activated Fe<sup>3+</sup>- ligand will spontaneously scavenge tumor abundant Fe<sup>3+</sup> at the site of enzyme activity forming a highly

stable Fe-complex, to localize and accumulate the signal enhancement groups (*e.g.* Gd-DOTA or Gd-PCTA) in tumor, revealing regional  $\beta$ -gluc activity, and verifying the location and magnitude of tumor to evaluate the gene therapy. Also, the formation of the Fe-complex will restrict motion of the Gd<sup>3+</sup> chelates, then enhancing additional relaxivity. **Figure 4** depicts the mechanism for detection of *lacZ* gene expression through Fe<sup>3+</sup>-trapped MRI contrast agent formation.



**Figure 4.** Detection of *lacZ* gene expression by  $\beta$ -gal activated *in situ* Fe<sup>3+</sup>-trapped MRI contrast agent formation.

Prostate-specific membrane antigen (PSMA) is a type II transmembrane glycoprotein with enzymatic activities: N-acetylated  $\alpha$ -linked L-amino dipeptidase (NAALADase) and  $\gamma$ -glutamyl carboxypeptidase (folate hydrolase).[97-99] Studies with the monoclonal antibodies have demonstrated that PSMA is the most well-established, highly restricted prostate cancer cell surface antigen, it is expressed at high density on the cell membrane of all prostate cancers.[100-102] The high prostate tissue specificity of PSMA has been identified as an ideal therapeutic and diagnostic target for prostate cancer, this potential was exemplified by the recent FDA approval of an <sup>111</sup>In-labeled PSMA monoclonal antibody (Prostascint<sup>®</sup>) for diagnostic imaging of prostate cancer.[103-110] Furthermore, phase I and II trials have begun using immunotherapy directed against PSMA.[106-108] By introducing  $\gamma$ -glutamate residue instead of D-galactose in the Figure 4, we intend to develop a novel class of PSMA responsive Gd(III)-based MRI approach specific for prostate cancers detection with high sensitivity (**Figure 5**).

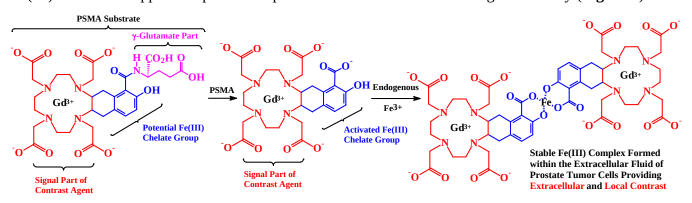


Figure 5. PSMA responsive Gd(III)-based MRI approach specific for prostate cancers detection.

Especially, PSMA has a large extracellular domain,[108] so the expression of PSMA tethered to the surface of the prostate cancer cells makes that the above novel peptide-based MRI contrast agents can be activated extracellularly around prostate cancers,[109] thus the need for a peptide-based MRI contrast agent to penetrate the prostate tumor cell membrane is no longer a prerequisite. The

permeability is always one of the greatest challenges in the development of *in vivo* MRI contrast agents.[111]

Accordingly, depending upon the enzyme sources either being the *lacZ* transgene or the PSMA from prostate tumors, this new platform could provide *in vivo lacZ* gene expression assay or *in vivo* prostate cancer imaging (in particular, through **extracellular** contrast agents), with combining the factors of reaching the high relaxivities. Furthermore, this new class of responsive MRI contrast agent is composed of three functional moieties, in which the signal enhancing and Fe<sup>3+</sup> chelating parts are changeable allowing modification in a search for ideal Fe<sup>3+</sup>-trapped MRI contrast agents. Importantly, the combination of three functional moieties is based on the clinically applied strategies on cancer therapy. These facts strongly suggest the potential of the proposal to future clinical application.

## STATEMENT OF WORK

**Specific Aim 1** Design and synthesize model "smart" MRI contrast agents to report  $\beta$ -gal or PSMA activities with the ability of trapping Fe<sup>3+</sup> ion.

Task 1 Design and optimization of synthetic strategies for reporter molecules. (Completed)

Task 2 Structural characterizations of the synthesized molecules. (Completed)

Specific Aim 2 Test the properties of molecules in solution and *in vitro* with cultured prostate cancer cells.

Task 3 Evaluation the basic properties of the agents in solution. (Completed)

Task 4 Evaluation of the properties of the optimal molecules *in vitro* with cultured prostate cancer cells. (Completed)

**Specific Aim 3** Scale up synthesis of the most promising MRI contrast agent(s) and apply to animal investigations.

Task 5 Scale up synthesis of the most promising <sup>1</sup>H MRI contrast agent(s). (Completed)

**Task 6** Apply the most promising <sup>1</sup>H MRI contrast agent(s) to assess  $\beta$ -gal transfection efficiency, *lacZ* gene expression (spatial and temporal) in prostate tumors *in vivo*. (**Completed**)

Task 7 Test dosing protocols, timing, MR detection protocols (Completed)

Task 8 Prepare manuscripts and final report (Completed)

# BODY

**SEYNTHESIS** Initially, we started the syntheses of the target molecules with the strategy of constructing the structures of  $Gd^{3+}$  and  $Fe^{3+}$  chelators simultaneously in the fused way as designed in the proposal, in order to maximize the restriction for the motion of the  $Gd^{3+}$  chelates, then obtaining the optimal relaxivity.

In the years 1 and 2, the syntheses according to the original plan met the challenges on (1) selective removal of benzyl ether (in blue) in the presence of benzyl ester (in magenta) by Pd/C hydrogenolysis; (2) selective removal of esters to accomplish the expected compound **D**, then to the target molecule  $M_1$  (see Figure 6). Because  $M_3 \sim M_6$  are analogues of  $M_1$  and  $M_2$ , similarly, their syntheses had encountered the same situations. Although we put in much time and effort for solving these issues even partially in year 3, the failure made us modify the synthetic strategy by using tert-Butyl (instead of Ethyl) (see Figure 7), since they can be readily and selectively removed.

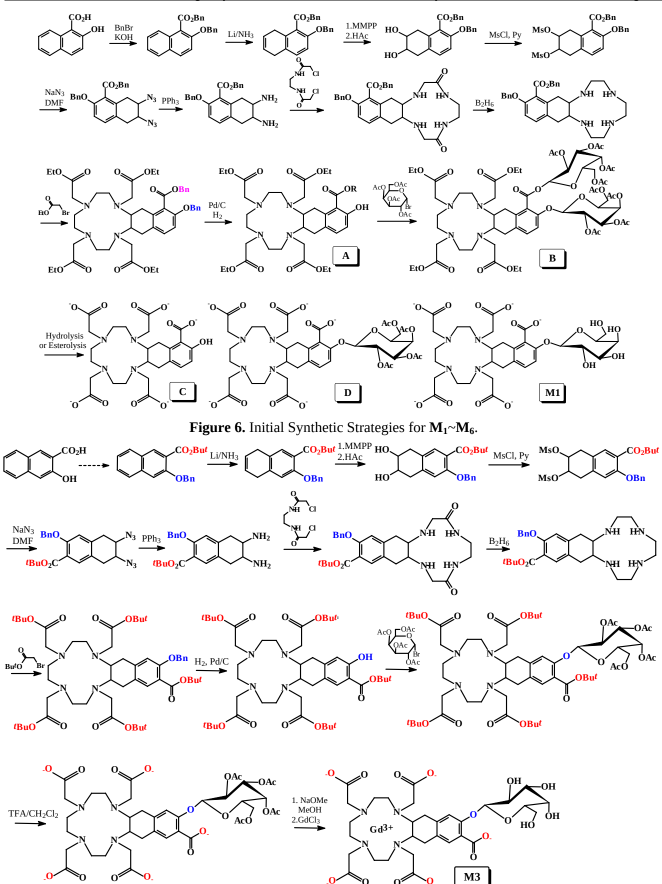
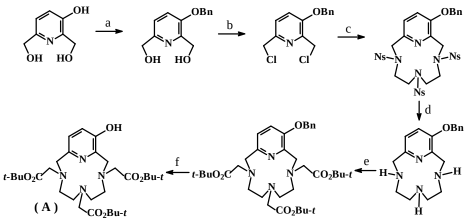
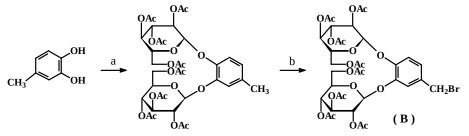


Figure 7. Modified Synthetic Strategies for M<sub>1</sub>~M<sub>6</sub>.

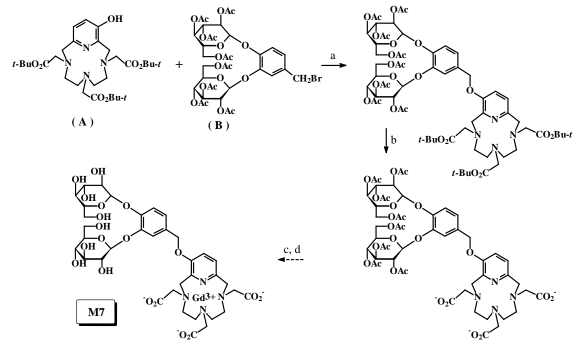
Following the modified synthetic strategy, we eventually achieved the designed target molecules  $M_1 \sim M_6$  and  $M_7$ ,  $M_8$  (see Figure 8).



**Reaction Conditions:** (a) BnBr, K2CO3, MeCN, 79%; (b) Bu3P, CCl4, 80%; (c) NsNH(CH2)2N(Ns)(CH2)2NHNs, K2CO3, MeCN, 81%; (d) HSCH2COOH, LiOH, DMF, 75%; (e) BrCH2CO2Bu-*t*, K2CO3, 85%; (f) H2, Pd/C, 90%.

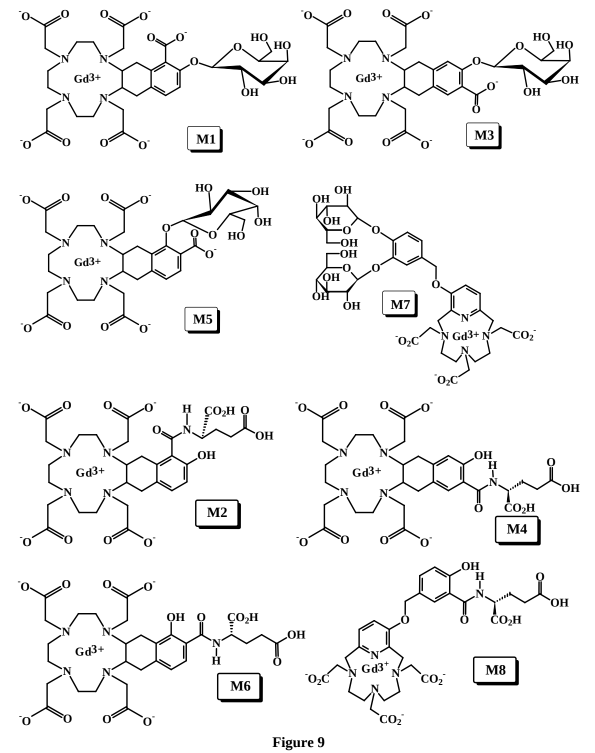


**Reaction Conditions:** (a) 2,3,4,6-tetra-*O*-acetyl-*a*-D-galactopyranosyl bromide, Hg(CN)<sub>2</sub>, 4Å M.S., CH<sub>2</sub>Cl<sub>2</sub>, 88%; (b) NBS, 78%;



**Figure 8. Reaction Conditions:** (a) K<sub>2</sub>CO<sub>3</sub>, MeCN, 75%; (b) CF<sub>3</sub>CO<sub>2</sub>H, CH<sub>2</sub>Cl<sub>2</sub>, 81%; (c) GdCl<sub>3</sub>, Pyridine, 82%; (d) MeOH, MeONa, 89%.

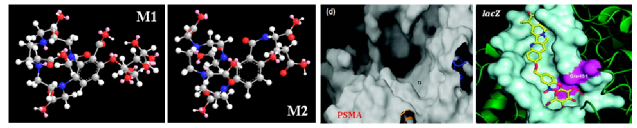
The test of the target molecules  $M_1 \sim M_8$  using <sup>1</sup>H-MRI by comparing the contrast enhancement with that of the control in sodium phosphate buffer solution (PBS) (0.1 M, pH=7.4) in the presence of ferric ammonia citrate (FAC) with  $\beta$ -galactosidase E801A or PSMA (from lysed LNCaP cells in Tris buffer) indicated that: (1) the reporter molecules  $M_1$ ,  $M_3$ ,  $M_5$ ,  $M_7$  can not be hydrolyzed by  $\beta$ -galactosidase E801A; (2) the reporter molecules  $M_2$ ,  $M_4$ ,  $M_6$ ,  $M_8$  cannot be hydrolyzed by PSMA; so no MRI contrast changes before and after addition of  $\beta$ -galactosidase E801A or PSMA can be seen (**Figure 9** and **Table** 1).



**Table 1.** <sup>1</sup>H-MRI contrast of the reporters  $M_1 \sim M_8$  in the presence of FAC. [Conditions: T<sub>1</sub>-weighted <sup>1</sup>H MRI, 200MHz, TR=300ms, TE=20ms, 1.5mm slice, 128×128, 50×50mm<sup>2</sup>. (A) Control,  $M_1$ ,  $M_3$ ,  $M_5$ ,  $M_7$  each (5µmol), FAC (2.5µmol), PBS (0.1M, pH=7.4, 1.5mL);  $M_2$ ,  $M_4$ ,  $M_6$ ,  $M_8$  each (5µmol), FAC (2.5µmol), Tris buffer (50 mM, pH=7.4, 1.5mL); (B) Complex,  $M_1 \sim M_8$  each (5µmol), FAC (2.5µmol), E801A (10 units), PBS (0.1M, pH=7.4, 1.5mL)], or PSMA (from lysed LNCaP cells 5×10<sup>6</sup>), Tris buffer (50 mM, pH=7.4, 1.5mL).

Enzyme	β-galactosidase E801A				PSMA				
Molecule	<b>M</b> <sub>1</sub>	<b>M</b> <sub>3</sub>	<b>M</b> <sub>5</sub>	$M_7$	$M_2$	$M_4$	<b>M</b> <sub>6</sub>	<b>M</b> <sub>8</sub>	
T <sub>1</sub> -weighted <sup>1</sup> H MRI (Control)							0		
T <sub>1</sub> -weighted <sup>1</sup> H MRI (Enzyme)									

The molecular modeling shows that the fused structure of  $Gd^{3+}$  and  $Fe^{3+}$  chelator results in the molecular structures of  $M_1 \sim M_8$  very rigid (*see* Figure 10 as an example of  $M_1$ ,  $M_2$ ). Therefore, we deduced that these too rigid molecules would be hard to coordinate with PSMA or *lacZ* proteins and dock into the cleft of the active site of the enzymes for interaction.[112,113]

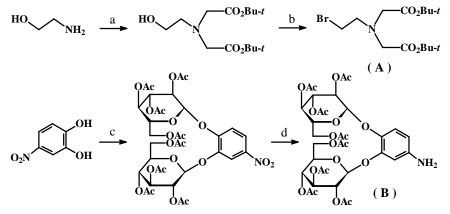


#### Figure 10

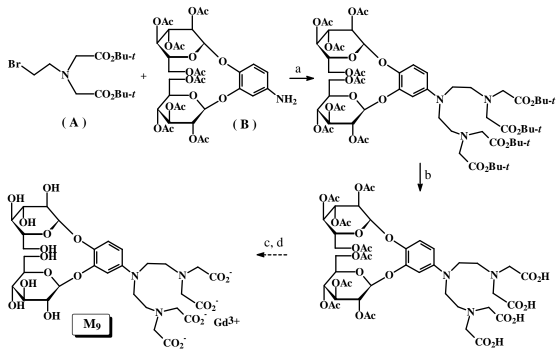
The molecular modeling and docking studies provide us a better understanding of the interactions between our designed target molecules and PSMA or *lacZ* proteins. We found that selecting a suitable structure with certain features (*e.g.* flexibility, linkage, appropriate angles, versatile binding modes, and coordination to  $Fe^{3+}$  with plastic chelating geometry) is crucial to the construction of the enzyme responsive enhanced MRI contrast agents. We also realized that the previous design and synthesis involved too many steps of reactions, not briefly and efficiently, displaying the increasing difficulty either in synthesis and purification with lower yields, especially, when the molecules grew bigger.

With these considerations in mind, we tried to introduce diethylenetriamine-N,N',N'',N'', tetraacetate (DTTA) instead of the cyclic DOTA or PCTA as Gd(III) chelator (see **Figure 11**) with a straightforward strategy. Most importantly, molecule **M**<sub>9</sub> can be hydrolyzed by  $\beta$ -galactosidase E801A in the presence of FAC in PBS (0.1 M, pH=7.4), producing obvious MRI contrast change before and after reaction with  $\beta$ -galactosidase E801A (see **Figure 12**), implying we found the right way. Inspirited by these results, we extended this strategy further for the synthesis of other new molecules by using the Fechelation agents in anticancer therapy as the Fe-chelator for construction of the responsive MRI contrast agents. We firstly proved that the clinically applied Fe-chelators, such as **Pyridoxal IsonicotinoylHydrazone (PIH)**, and its analogues **Salicylaldehyde BenzoylHydrazone (SNH)**, can act as

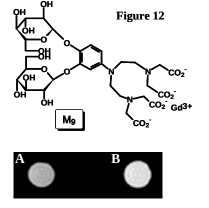
Fe-based <sup>1</sup>H MRI contrast agents to produce strong  $T_1$ -weighted contrast effects (**Figure 13**, **TABLE 2**), suggesting that the Fe-complex formation could not only localize, accumulate and restrict the motion of the linked Gd-based <sup>1</sup>H MRI moiety, but also itself can produce the additional relaxivity.



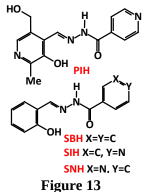
**Reaction Conditions:** (a) BrCH<sub>2</sub>CO<sub>2</sub>Bu-*t*, KHCO<sub>3</sub>, 88%; (b) Ph<sub>3</sub>P, NBS, 86%; (c) 2,3,4,6-tetra-*O*-acetyl-*a*-D-galactopyranosyl bromide, Hg(CN)<sub>2</sub>, 4Å M.S., CH<sub>2</sub>Cl<sub>2</sub>, 92%; (d) H<sub>2</sub>, Pd/C, 100%.



**Figure 11. Reaction Conditions:** (a) K<sub>2</sub>CO<sub>3</sub>, MeCN, 78%; (b) CF<sub>3</sub>CO<sub>2</sub>H, CH<sub>2</sub>Cl<sub>2</sub>, 84%; (c) GdCl<sub>3</sub>, Pyridine, 80%; (d) MeOH, MeONa, 86%.



**Figure 12**.  $T_1$ -weighted (TR/TE 250/12 ms) MR images of solutions and the signal intensity in test tubes at 4.7 T MR scanner: (**A**) PBS with **M9** and FAC; (**B**) PBS with **M9**, FAC and βgalactosidase E801A.

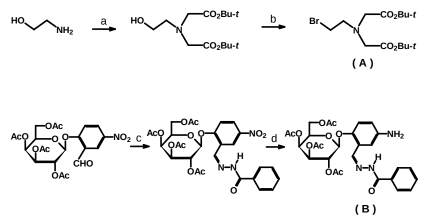


Then, we synthesized  $M_{10}$ ,  $M_{11}$  and  $M_{12}$ by using SBH, SIH and SNH as the Fe-chelators (see Figure 14). The MRI evaluation of the reporter molecules  $M_{10}$ ,  $M_{11}$  and  $M_{12}$ , respectively, in sodium phosphate buffer solution (PBS) (0.1 M, pH=7.4) in the presence of ferric ammonia citrate (FAC) with  $\beta$ galactosidase E801A indicated that: (1) Again, the reporter molecule  $M_{10}$ , similarly like  $M_9$ , can be hydrolyzed by  $\beta$ -galactosidase E801A,

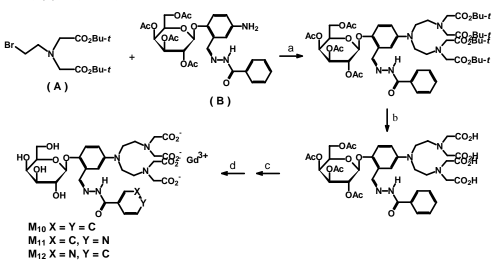
Chelator	PIH	SBH	SIH	SNH
$T_1$ -weighted <sup>1</sup> H MRI (Control) <sup>1</sup>				
$T_1$ -weighted <sup>1</sup> H MRI (Control) <sup>2</sup>				
$T_1$ -weighted <sup>1</sup> H MRI (Complex)				

**TABLE 2.** (A) (1) Control, PIH, SBH, SIH or SNH each (1.6mM) in PBS; (2) Control, ferric ammonia citrate (FAC) (0.8mM) in PBS; (B) Complex, PIH, SBH, SIH or SNH each (1.6mM), FAC (0.8mM) in PBS; **CONDITIONS**: T<sub>1</sub>-weighted <sup>1</sup>H MRI, 200MHz, TR=300ms, TE =20ms, 1.5mm slice, 128×128, 50×50mm<sup>2</sup>.

producing apparent MRI contrast change upon response to  $\beta$ -galactosidase E801A; (2) Unlike M<sub>9</sub> and M<sub>10</sub>, M<sub>11</sub> and M<sub>12</sub> have no MRI contrast enhancements with galactosidase E801A in the presence of FAC in PBS (0.1 M, pH=7.4), but with strong MRI contrast changes with galactosidase G5160 in the presence of FAC in PBS (0.1 M) at pH=4.5 (**Table 3**).



**Reaction Conditions:** (a) BrCH<sub>2</sub>CO<sub>2</sub>Bu-*t*, KHCO<sub>3</sub>, 88%; (b) Ph<sub>3</sub>P, NBS, 86%; (c) PhCONHNH<sub>2</sub>, EtOH, AcOH, Reflux, 82%; (d) H<sub>2</sub>, Pd/C, 96%.



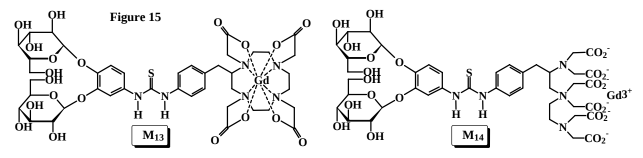
**Figure 14. Reaction Conditions:** (a) K<sub>2</sub>CO<sub>3</sub>, MeCN, 42%; (b) CF<sub>3</sub>CO<sub>2</sub>H, CH<sub>2</sub>Cl<sub>2</sub>, 64%; (c) GdCl<sub>3</sub>, Pyridine, 70%; (d) MeOH, MeONa, 85%.

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**Table 3**. <sup>1</sup>H-MRI contrast of the reporters  $M_{10} \sim M_{12}$  in the presence of FAC with E801A in PBS (0.1M, pH=7.4) or G5160 in PBS (0.1M, pH=4.5). [**Conditions:** T<sub>1</sub>-weighted <sup>1</sup>H MRI, 200MHz, TR=300ms, TE=20ms, 1.5mm slice, 128×128, 50×50mm<sup>2</sup>. (A) Control,  $M_{10} \sim M_{12}$  each (5µmol), FAC (2.5µmol), PBS (0.1M, pH=7.4, 1.5mL); (B) Complex,  $M_{10} \sim M_{12}$  each (5µmol), FAC (2.5µmol), E801A (10 units), PBS (0.1M, pH=7.4, 1.5mL)], or G5160 (10 units), PBS (0.1M, pH=4.5, 1.5mL)].

Enzyme	β-galactosidase E801A			β-galactosidase G5160			
Molecule	$M_{10}$	M <sub>11</sub>	<b>M</b> <sub>12</sub>	$M_{10}$	$M_{11}$	M <sub>12</sub>	
T <sub>1</sub> -weighted <sup>1</sup> H MRI (Control)							
T <sub>1</sub> -weighted <sup>1</sup> H MRI (Enzyme)							

The experience accumulated on the development of enzyme responsive enhanced MRI contrast agents opened our mind, and the desire for an ideal *in vivo* MRI probe prompted us to design and syntheses another two kinds MRI agents: (1) through phenylthioureido as linkage connecting  $Gd^{3+}$  and  $Fe^{3+}$  chelators for suitable flexibility of the molecules  $M_{13}$  and  $M_{14}$  (see Figure 15), both like  $M_9$  produced apparent MRI contrast differences upon response to  $\beta$ -galactosidase E801A.



(2) "Click Chemistry" Approach Because of regioselectivity, high yields in reasonable reaction times under mild conditions, "Click Chemistry" has been applied in a wide range of fields from synthetic chemistry to biomedicine and materials science. Our attention is on the versatile triazole rings as linkers between Fe<sup>3+</sup> and Gd<sup>3+</sup>-ligands to functionalize with tolerance for the interaction with *lacZ* protein (see **Figure 16**).

*In Vitro* MRI Studies (1) Cell preparation (a) Stably transfected PC3 cell line: *E. coli lacZ* gene (from pSV- $\beta$ -gal vector, Promega, Madison,WI) was inserted into high expression human cytomegalovirus (CMV) immediate-early enhancer/promoter vector phCMV (Gene Therapy Systems, San Diego, CA) giving a recombinant vector phCMV/*lacZ*, which was used to transfect PC3 cells using GenePORTER2 (Gene Therapy Systems). Cells were grown in DMEM (Dulbecco's Modification of Eagle's Medium, Mediatech, Inc, Herndon, VA), 10% FBS (Fetal bovine serum, Hyclone, Logan, UT) with 1% Penicillin-streptomycin Solution (Mediatech). The highest  $\beta$ -gal expressing colony was selected using G-418 disulfate (C<sub>20</sub>H<sub>40</sub>N<sub>4</sub>O<sub>10</sub>. 2H<sub>2</sub>SO<sub>4</sub>, Research Products International Corp, Mt. Prospect, IL) (800 µg/ml), which was also included for routine culture (200 µg/ml). with 1% Penicillin-streptomycin Solution (Mediatech). The highest  $\beta$ -gal expressing G-418 disulfate

 $(C_{20}H_{40}N_4O_{10}. 2H_2 SO_4, Research Products International Corp, Mt. Prospect, IL) (800 µg/ml), which was also included for routine culture (200 µg/ml).$ 

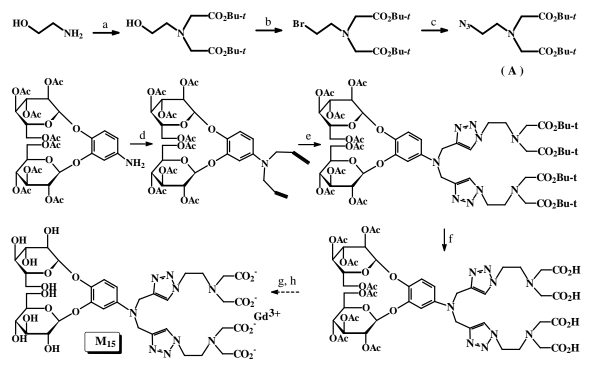
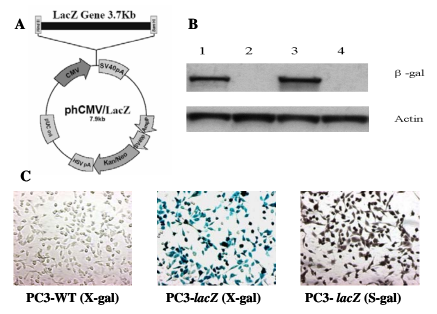


Figure 16. Reaction Conditions: (a) BrCH<sub>2</sub>CO<sub>2</sub>Bu-*t*, KHCO<sub>3</sub>, 88%; (b) Ph<sub>3</sub>P, NBS, 86%; (c) NaN<sub>3</sub>, DMF, 80C, 88%, (d) CHCCH<sub>2</sub>Br, K<sub>2</sub>CO<sub>3</sub>, DMF, rt, 86%, (e) (A), CuSO<sub>4</sub>, NaAsc, t-BuOH, rt, 69%, (f) CF<sub>3</sub>CO<sub>2</sub>H,

CH<sub>2</sub>Cl<sub>2</sub>, 74%; (g) GdCl<sub>3</sub>, Pyridine, 71%; (h) MeOH, MeONa, 82%.

X-gal **(b)** and S-gal staining for  $\beta$ -gal: cells were PBS plus fixed in 0.5% glutaraldehyde (5 min) and rinsed in PBS prior to staining. Staining was performed using standard procedures for 2 hours at 37 °C in PBS plus 1 mg/ml Xgal (Sigma, St. Louis, MO), 1 mM MgCl<sub>2</sub>, 5 mM K<sub>3</sub>Fe(CN)<sub>6</sub>. and 5 mM K<sub>4</sub>Fe(CN)<sub>6</sub> or with 1.5 mg/ml S-gal (Sigma) and 2.5 mg/ml FAC (see Figure 17). (c)  $\beta$ -Gal activity assay: The  $\beta$ -gal activity of tumor cells and tissues in mice was measured using the β-gal assay kit (Promega, Madison, WI) with yellow o-



**Figure 17** Generation of PC3 cells stably expressing of  $\beta$ -gal. (**A**) Map of recombinant *lacZ* vector (phCMV/*lacZ*). (**B**) Western blot: cell extracts of two transfected lines PC3-*lacZ*1 (lane 1) and PC3-*lacZ* (lane 3), together with PC3-WT (lanes 2 and 4) were examined. (**C**) PC3 wild-type and PC3-*lacZ* cells were stained using X-gal and S-gal: over 90% of PC3-*lacZ* cells were stained blue or black, respectively, while the PC3 wild type cells did not stain.

nitrophenyl  $\beta$ -D-galactopyranoside. (d) Western blot analysis: Protein was extracted from PC3 tumor cells and was quantified by a protein assay (Bio-Rad, Hercules, CA) based on the Bradford method. Each well was loaded with 30µg protein and separated by 10% SDS-PAGE (Nu-PAGE) and transferred to a polyvinylidene fluoride (PVDF) membrane. Primary monoclonal anti- $\beta$ -gal antibody (Promega) and anti-actin antibody (Sigma) were used as probes at a dilution of 1:5000, and reacting protein was detected using a horseradish peroxidase-conjugated secondary antibody and ECL detection (Amersham, Piscataway, NJ).

(2) In Vitro MRI The in vitro evaluation of  $M_{9}$ ~ $M_{14}$  with PC3-lacZ cells in the presence of FAC showed that only molecules  $M_9$  and  $M_{10}$  exhibited apparent MRI differences.  $M_9$  and  $M_{10}$  (6 µmol) each in 1:1 DMSO/PBS was added to suspensions of  $5 \times 10^6$  PC3 wild type and PC3-lacZ cells in PBS (1.0 mL) and FAC (3 µmmol) in wells and maintained at 37°C. MRI experiments were performed on a 4.7 T Varian Unity INOVA spectrometer. Figure

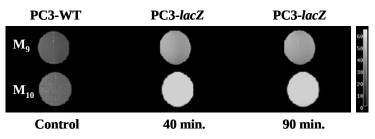
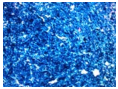


Figure 18 <sup>1</sup>H MRI, 200 MHz, TR=300ms, TE=20ms, 1.5mm slice,  $128 \times 64$ ,  $40 \times 40$  mm<sup>2</sup>. (A) control, M<sub>9</sub> or M<sub>10</sub> (6 µmol), FAC (3 µmol), 5×106 PC3 WT, PBS (0.9 mL), DMSO (0.1 mL); (B) M<sub>9</sub> or M<sub>10</sub> (6 µmol), FAC (3 µmol), 5×106 PC3-lacZ, PBS (0.9 mL), DMSO (0.1 mL)].

18 showed the *in vitro* MR images of  $M_9$  and  $M_{10}$  with *lacZ* transfected prostate tumor cells, yielding obvious MRI contrast changes between in WT and lacZ transfected PC3 prostate tumor cells, indicating that both M<sub>9</sub> and M<sub>10</sub> can penetrate prostate tumor PC3 cell membrane and have no apparent cytotoxicity and no physiological perturbation effects on WT and lacZ transfected PC3 cells, the others M<sub>11</sub>~M<sub>14</sub> cannot cross prostate tumor PC3 cell membrane.

In Vivo MRI Studies of  $M_9$  and  $M_{10}$  (1) Animal model All in vivo MRI studies were performed with approval from the Institutional Animal Care and Use Committee (IACUC). Wild type and stably transfected *lacZ* PC3 cells  $(2 \times 10^6)$  were implanted subcutaneously in the left and right thighs of mice, respectively, when the tumors reached  $\sim 0.8$  cm in diameter, the mouse was anesthetized (1.3% isoflurane/air at 1 dm<sup>3</sup>/min) with a facemask and maintained at 37°C by a warm pad with circulating water, and placed into animal coil for imaging. MRI data were acquired using a 4.7 T horizontal bore magnet with a Varian INOVA Unity system (Palo Alto, CA, USA).  $T_1$  and  $T_2$  values were measured using a spin echo sequence with varying repetition and echo times, e.g. T<sub>1</sub>-weighted <sup>1</sup>H MRI, 200MHz, TR=0.2, 0.3, 0.5, 0.8, 1, 1.5, 2, 4, 6s,





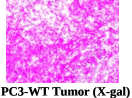


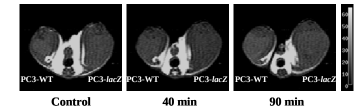
Figure 19

TE=12ms, 1.5mm slice, matrix=128×128, FOV=50×50mm<sup>2</sup>;  $T_2$ -weighted <sup>1</sup>H MRI, 200MHz, TR=6s, TE=11, 15, 20, 30, 50, 100, 150ms, 1.5mm slice, matrix=128×128, FOV=50×50mm<sup>2</sup>. Histology analysis confirmed that PC3-lacZ tumor section showed over 90% of tissue stained blue for  $\beta$ -gal, while PC3-WT tumor histological section showed little or no blue stain (Figure 19). (2) In Vivo MRI with i.v. injection Mice bearing PC3-WT and PC3-lacZ tumors were imaged on a 4.7 T Varian Unity INOVA spectrometer. T1-weighted transaxial images were obtained before and after intravenous injection of the W81XWH-05-1-0593 - Final Report Yu, Jian-Xin 16

mixture of 0.4 mmol/kg M<sub>9</sub> and M<sub>10</sub> and FAC. Postcontrast scans were obtained every 15 min for one

and half hours. For both reporters  $M_9$  and  $M_{10}$ , the MR images of animals showed that there are no time-signal intensity changes between PC3-WT and PC3-*lacZ* tumors before and after  $M_9$  and  $M_{10}$  injection (Figure 20), indicating that both  $M_9$  and  $M_{10}$  can either be washed out or metabolized very quickly, and can't reach to PC3-WT and PC3*lacZ* tumors on the thighs with enough

amount. Also, we found that mice all died one and half-hours later after intravenous injection of M<sub>9</sub>. (3) *In Vivo* MRI with direct injection into tumors However, if a solution of M<sub>10</sub> (0.4 mmol/kg) and FAC (DMSO/PBS 1:1 V/V') was injected directly into the tumors



**Figure 20** <sup>1</sup>H MRI, 200 MHz, TR=250ms, TE=12ms, 1.5mm slice, 128×64, 40×40 mm<sup>2</sup>. (A) control; (B) M<sub>10</sub> (0.4 mmol/kg), FAC (0.2 mmol/kg), PBS/DMSO (1:1), i.v. injection].

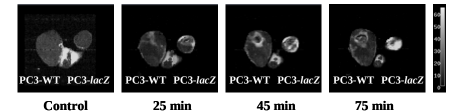


Figure 21 <sup>1</sup>H MRI, 200 MHz, TR=250ms, TE=12ms, 1.5mm slice, 128×64, 40×40 mm<sup>2</sup>. (A) control; (B)  $M_{10}$  (0.4 mmol/kg), FAC (0.2 mmol/kg), PBS/DMSO (1:1), i.v. injection].

in a "fan" pattern, strong contrast was detected in the *lacZ* expressing PC3 tumors (Figure 21).

# **RESEARCH ACCOMPLISHMENTS**

(1) Designed and synthesized a series of reporter molecules, and verified their structures, importantly, accumulated solid foundation, experience and expertise for the further investigation on molecular imaging.

(2) Finished the *in vitro* and *in vivo* evaluation of the reporter molecules  $M_9$  and  $M_{10}$ , and the results demonstrated this novel mechanism proposed in W81XWH-05-1-0593 for imaging and evaluation of prostate cancer gene therapy is reliable.

#### **REPORTABLE OUTCOMES**

(1) A series of abstracts had been accepted for presentation on the various conferences such as World Molecular Imaging Congress, *Innovative Minds in Prostate Cancer Today*, American Chemical Society Meeting.

(2) Several papers are in preparation.

#### **CONCLUSIONS**

Prostate cancer is the most commonly diagnosed cancer and the second most common cause of cancer death in men in the United States. The advent of effective screening measures can sharply decrease the mortality of prostate cancer through detecting this disease at an earlier stage. However, the evidence for mortality benefit from prostate cancer screening has been disappointing to date. Expanding knowledge of prostate cancer biology with combination of imaging technologies would be of considerable value in many ongoing and future clinical prostate cancer diagnosis and gene therapy trials. W81XWH-05-1-0593 – Final Report 17 Yu, Jian-Xin

Based on the biologic features of prostate cancer, we proposed in this project a new approach for *in vivo lacZ* gene expression assay or *in vivo* prostate cancer imaging. The ultimate objective is to demonstrate the utility and reliability of this new approach to measure  $\beta$ -gal or PSMA activities *in vivo*. We have accomplished a series of target molecules  $M_1 \sim M_{14}$ , and verified by NMR data. Strong MRI contrast changes of target molecules  $M_9$  and  $M_{10}$  for detection *lacZ in vitro* and *in vivo* demonstrated this novel mechanism described in W81XWH-05-1-0593 is feasible and reliable. With screening out the ideal reporter molecules, we believe the translation of this novel approach to clinical investigations will enable prostate cancer detection comprehensive and infallible.

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# DEVELOPMENT OF NOVEL Fe-BASED <sup>1</sup>H MRI lacZ GENE REPORTERS FOR IN VIVO ASSESSMENT

# **JT SOUTHWESTERN** MEDICAL CENTER

Jian-Xin Yu, Dawen Zhao, Ralph P. Mason

Cancer Imaging Program, Department of Radiology

Presentation: 0614

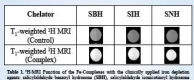
# University of Texas Southwestern Medical Center at Dallas, Texas, USA

MOLECULAR DESIGN

INTRODUCTION The lacZ gene encoding β-galactosidase (β-gay) attractive reporter gene, and its introduction has become a trave-structive reporter gene, and its introduction has become a trave-tic nonirvative in orice detection we be an original cancer gene sidase (β-gal) has been recognized as the mos ntroduction has become a standard means of sion, and uld be of onsiderable value in many ongoing and future clinical cancer gene therapy rials.<sup>01</sup>To this end, a variety of *in vivo lacZ* gene reporters has been developed vivo lacZ gene reporters has be nt.<sup>[2-5]</sup> bioluminescent.<sup>[6]</sup> rac atric A fh ent @ radiot such as colorimetric,<sup>14</sup> Biorescent,<sup>24</sup> bioluminescent,<sup>24</sup> radiotracers for positron emission tomography (PET) or alingle-photon emission computed tomography (SPECT))<sup>174</sup> magnetic resonance imaging (MRI) probes<sup>30,11</sup> and <sup>17</sup> magnetic resonance spectroscopy (MRS) and chemical shift imaging (CS) reporters:<sup>124</sup> We now report the development of a new class of Fe-based <sup>14</sup> MRI *lacZ* gene reporters based on clinically applied iron-depletion

tically important metal ion for a wide variety of cellular e as compared with their normal counterparts, frequently et Tumor cells, as compared with their normal counterparts, frequently exhibit increased uptake and utilization of more iron. Many studies indicate that a high level of iron accumulated in animals and humans is associated with both Vet or non-accumulate an annihals and numbran is astocated with ooth ation and the progression of cancers. Cancer cells are also sensitive depletion because of the critical requirement for iron in proteins that entitl a roles in DNA synthesis, repair and energy production. Iron on as a therapeutic strategy has been applied for cancers. The FDA has approved five iron cheators for use in anticancer therapy, others are valuated in clinical settings.<sup>840</sup>

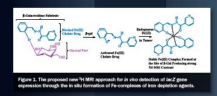
Based on the structural similarities between the iron complexes with the clinically applied iron depletion agents and the known Fe-based 'IH MRI contrast agents, we firstly demonstrated that the clinically applied Fe-chelators can act as Fe-based 'IH MRI contrast agents producing strong contrast effects (Table 1), opening a new transition of the order on agent as anticancer drug as well as of 'IH MRI detection or diagnosis agent.



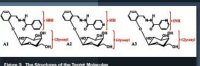
(SIH) and its analog salicylaldehyde ni (SIH) and its analog salicylaldehyde ni 'H MRI, 200MHz, TR=300ms, TE=2 FAC (2.5µmol), PBS (1.5mL); (B) (2.5µmol), PBS (1.5mL)]. icotinoyi hydrzzone (SNH). [Conditions: Ti-wa icotinoyi hydrzzone (SNH). [Conditions: Ti-wa Ima, 1.Jmm alice, 128-128, 50-30mm<sup>2</sup>, (A) Ce Complex, SBH, SIH or SBH auch (5µmol),

#### MOLECULAR DESIGN

Prompted by this finding, we now propose a novel molecular design to deliver the in situ formation of Pe-complexes of iron depletion agents specific targeting on cancer by using Gene Directed Enzyme Prodrug Therapy (GDEPT)

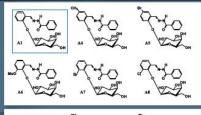


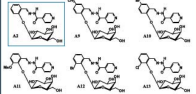
#### TARGET MOLECULE SYNTHESIS

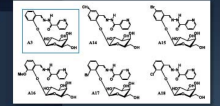


#### ALTERNATE MOLECULE SYNTHESIS

lering the multiple requirements for an ideal *in vivo* reporter, v ed a series of alternate candidates based on the different substituents desi diff ed a series of alternate candidates based on the different substituents at nt positions, which shall be directly related to the enzyme sensitivity ecificity, aqueous solubility, toxicity, and importantly, the ability of the d arcythydrazones to complex Fe+ ion (Fe-based 'H MRI effects).







#### RESULTS

Testing A1-A18 with  $\beta$ -galactosidase G-5160, showed that all, except A7, A8, A12 and A13, can react with  $\beta$ -galactosidase G-5160 in the presence of Fe<sup>3</sup>. The released iron depletion agents then spontaneously capture Fe<sup>3</sup> readily in situ forming stable Fe-complexes, which produce strong Fe-based <sup>1</sup>H MRI contrast.

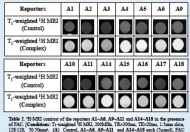


Table 2. 'B-MRI contrast of the reporter: Al-A6, A9-A11 and A14-A15 in the pre-of FAC. [Conditions: Tr-weighted H MRI, 2000;Hz, TR=300m, TR=20m, 1. Junn 128126, 59. 'Source', (A). Control, Al-A6, A9-A11 and A14-A18 set (Symon), (2. Spund), PBS (0.1M, pH=-5, 1. Junl); (B). Complex, A1-A6, A9-A11 and A14 set (Symon), FAC (Symon), G-S100 (Sum), PBS (D). (M), pH=54, 1. Smal).]

#### CONCLUSIONS

These molecules demonstrate the feasibility of combining anticancer drug iron depletion agents for detection of the lacZ reporter gene, opening a new possibility of merging cancer therapy and detection into one approach.

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Dort ME et al., Mol Imaging, 2008, 7, 187 • AY et al. Annew. Chem. Inti. Edn. Engl. njugate Ch

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# DEVELOPMENT OF NOVEL lacZ RESPONSIVE ENHANCED **GD-BASED MRI AGENT**



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Presentation: 0615

Cancer Imaging Program, Department of Radiology

University of Texas Southwestern Medical Center at Dallas, Texas, USA

SYNTHESIS

STRUCTURAL FEATURES

#### INTRODUCTION

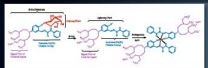
Gene therapy holds great promise for treating cancer an exploited in several clinical trials. A major current obst is to establish a method of assessment of therapeu of heterogeneity and longevity in tissues tosidase (β-gal) has been recognized as t herefore, noninyasive detection *in vivo* wor nized as the m β-galactoidase (β-gal) has been recognized as the most attraction gene. Therefore, noninvasive detection *in vivo* would be of comit in future elinical cancer gene therapy trials.<sup>(1)</sup> various groups has innovative approaches to assessing β-gal activity *in vivo* using enhanced contrast <sup>1</sup>H MRI or <sup>10</sup>F NMR spectroscopy.<sup>PA o</sup> or acionuclide imaging.<sup>(1)</sup> VM wor report the exploration of a ne to *lacZ* responsive Gd-based <sup>1</sup>H MRI agents.

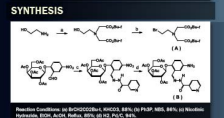
Descent and demonstrated that, by chelating Gd(phen)HDO3A with Fe(II) to form a highly stable tria-complex, the relaxivity increased 145% at 200HHz and 37°C(B<sup>3</sup>) Most recently. Toth dr al howed that the hearementalitic,self-assembled metallocar with itx efficiently relaxing GdIII centers, $[Fe(Gd_L(H,O)_1)<sup>4</sup>, exhibited a particularly high relaxivity for its moderate$ molecular weight.<sup>100</sup>

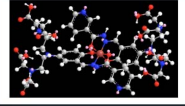
on is a critically important metal ion for a wide variety of cellular events immor cells, as compared with their normal counterparts, frequently exhibit reased uptake and utilization of more iron. Many studies indicate that high level of iron accumulation in animals and humans is associated a tugh repert of invitation and the progression of cancers. Cancer cells are with both the initiation and the progression of cancers. Cancer cells are also sensitive to iron depletion because of the critical requirement for iron in proteins that play essential roles in DNA synthesis, repeats and energy production. Iron depletion as a therapeutic strategy has been applied for cancers. The FDA has already approved five iron chelators for use in anticancer therapy, others are being evaluated in clinical settings.<sup>(9)</sup> energy ed for

#### DESIGN

pred by this finding, we now propose a novel class of enzyme activated based MRI contrast agent for *in vivo* detection of β-gal activity combing formes: (A) a signal enhancement group, such as Gd-DTTA; (B) and rapy vage by  $\beta$ -gal in cells, the r ie Fe<sup>3</sup> ligand will ming a his hly stable com

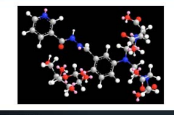




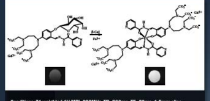


Fe-C

Ligand St



RESULTS



#### CONCLUSIONS

We propose a novel class of enzyme activated Gd<sup>36</sup>-based MRI contrast age for detection of  $\beta$ -gal activity. Synthesis of the substrate has been successf and the evaluation of this agent with  $\beta$ -gal aboved that the released aglyco including the activated Pe<sup>3</sup>-ligand and MRI signal enhancement gro spontaneously traps Pe<sup>36</sup> in the solation forming a highly stable complex, th retricting the motion of the Gd<sup>36</sup> chelates enhancing relaxity. We are n g this agent with  $\beta$ -gal expressing cells and in view

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ADDITIONAL INFORMATION For additional informa ion, p

#### -Xin VietterSo

