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[¹¹C]Ascorbic and [¹¹C]Dehydroascorbic Acid, An Endogenous Redox Pair for Sensing Reactive Oxygen Species Using Positron Emission Tomography

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Abstract

Here we report the radiosynthesis of an endogenous redox pair, $[^{11}C]$ ascorbic acid ($[^{11}C]$ VitC) and $[^{11}C]$ dehydroascorbic acid ($[^{11}C]$ DHA), the reduced and oxidized forms of vitamin C, and their application to ROS sensing. These results provide the basis for *in vivo* detection of ROS using positron emission tomography (PET).

Reactive oxygen species (ROS) are generated as a normal product of oxidative metabolism and are required signalling molecules in a diverse array of biological processes.¹ Dysregulation of ROS in common disease states including cancer,² neurodegeneration,³ chronic inflammation,⁴ and diabetes⁵ provides a powerful motivation to develop non-invasive biomarkers of oxidative stress. Current ROS sensing techniques in living systems are largely limited to *in vitro* study. Advances toward *in vivo* ROS detection include approaches using electron spin trapping (ESR),⁶ near-IR optical,⁷ bioluminescent,⁸ [¹³C] magnetic resonance imaging (MRI),^{9,21} chemiluminescent probes,¹⁰ fluorescent probes¹¹ and positron emission tomography (PET).¹² Due to its high sensitivity, good spatial resolution and low toxicity,¹³ PET has potential for detecting ROS in a clinical setting.

VitC is transported into cells via the sodium dependent vitamin C transporter (SVCT1–2).¹⁴ In the presence of ROS, VitC undergoes a two-electron oxidation to DHA, which in aqueous solution exists predominantly in bicyclic hemiketal form and is a substrate for glucose transport (GLUT 1, 3, 4).¹⁵ We hypothesized that by taking advantage of rapid GLUT

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transport,¹⁶ this mechanism could be employed to detect extracellular ROS (Figure 1). Previously Yamamoto¹⁷ and Kothari¹⁸ have investigated 6-[¹⁸F]-fluoro-6-deoxy-L-ascorbic acid as a PET analogue of VitC; however this probe cannot enter cells via GLUT, as the ¹⁸F label in the 6 position prevents formation of the bicyclic species of DHA.^{17b,17e,f} We have therefore developed a new pair of endogenous PET radiotracers [¹¹C] ascorbic acid ([¹¹C]VitC) and its oxidized partner, [¹¹C] dehydroascorbic acid ([¹¹C]DHA) and have used this redox pair to sense ROS *in vitro*.

[¹¹C]VitC was synthesized from L-xylosone based on a modification of the previously reported [^{13/14}C] enriched techniques (Scheme 1).¹⁹ The methods employed and relevant analytical data are reported in full in the Supporting Information (Figures S1-S3). Table 1 summarizes radiochemical yields for synthesis with varying amounts of added KCN carrier. With no carrier added [¹¹C]VitC in situ oxidation to [¹¹C]DHA was observed at pH 7 (Figure S4) possibly related to generation of ROS by radiolysis. This phenomenon has been previously observed for 6-[¹⁸F]-Fluoro-6-deoxy-L-ascorbic acid.¹⁸ Thus, we used nonradioactive carrier VitC to protect against in situ oxidation. This was achieved by adding carrier KCN during the radiochemical preparation. With the presence of 0.6 - 1.0 mM(specific activity $\approx 3.0-10.0$ mCi/µmol; 110–370 MBq) carrier in the final isolated product $[^{11}C]$ VitC is stable at all time points tested (Figure S5). As sampling of our institution's clinical 2-deoxy-2-[¹⁸F]fluoroglucose ([¹⁸F]FDG) doses revealed that the administered [¹⁸F]FDG solution contained 1.2 ± 0.1 mM (n = 3) non-radioactive glucose, we do not anticipate that addition of carrier at this level will significantly diminish the ability to image ^{[11}C]DHA transport via GLUT. Other antioxidants were also considered to prevent ^{[11}C]VitC *in situ* oxidation, but given that these also react with ROS they would likely confound interpretation of in vitro and in vivo data.

We first evaluated the transport of both [¹¹C]VitC and [¹¹C]DHA in U87 human glioblastoma cells using [¹⁸F]FDG as a standard radiotracer for GLUT transport. GLUT blocking studies were carried out with application of 10µg/ml cytochalasin B, a potent inhibitor of GLUT transport.^{15a,23} SVCT transport was interrogated by modulating (+)/(–) co-transport of Na⁺ as per Vera et al.^{15b–d} While uptake of [¹¹C]VitC is not affected by blocking of the GLUT receptor, uptake is notably decreased in the absence of Na⁺ (Figure 2a). However, uptake of [¹¹C]DHA is not affected by availability of Na⁺ for co-transport, and is effectively blocked by application of cytochalasin B (Figure 2b) mirroring the trend observed for [¹⁸F]FDG (Figure 2c). This data confirms previously reported trends for uptake of the non-radioactive compounds.¹⁵ Since the uptake of [¹¹C]DHA via GLUT is 10 fold higher than uptake of [¹¹C]VitC we anticipated that intracellular accumulation of [¹¹C]VitC via SVCT occurs more slowly than transport of the oxidized species, [¹¹C]DHA via GLUT in many tissues.

Having shown the expected behavior of $[^{11}C]$ VitC and $[^{11}C]$ DHA *in vitro*, we performed a proof of concept *in vivo* experiment to demonstrate the differential transport of the two tracers. It has been well-established that DHA (but not VitC) crosses the blood-brain barrier transported primarily by GLUT1.²³ Approximately 200 µCi of $[^{11}C]$ VitC (n = 3) and $[^{11}C]$ DHA (n = 3) each were administered to normal rats via tail vein injection and a 40 min

dynamic scan was obtained using a microPET/CT scanner. As anticipated the brain accumulation of $[^{11}C]$ DHA was markedly higher than that of $[^{11}C]$ VitC (Figure 3), confirming our hypothesis that changes in uptake based on oxidized vs. reduced forms of ascorbic acid can be detected using PET.

Finally we investigated ROS-dependent [¹¹C]VitC accumulation in cells. This was first accomplished in U87 cells by addition of exogenous H₂O₂ to the media,^{12a} resulting in a greater than 2-fold increase in $[^{11}C]$ accumulation (*p = 0.0006) as shown in Figure 4a. We next applied [¹¹C]VitC to a model of endogenous ROS production, namely stimulated neutrophil-lineage cells undergoing oxidative burst. For this study we used the HL60 cellline, a human leukemia neutrophilic precursor, and human neutrophils, which had been freshly isolated from whole blood. The mechanism of VitC uptake in human neutrophils has been well established in literature.^{24,15c} During phagocytosis, neutrophils undergo Noxmediated generation of ROS to destroy bacteria and simultaneously oxidize extracellular VitC.²⁵ [¹¹C]VitC was oxidized to [¹¹C]DHA by the major ROS produced during the oxidative burst, H_2O_2 , O_2^- and ClO⁻ (Figure S6). To investigate tracer uptake via this mechanism, cells were incubated with 10 μ Ci (0.37 MBq) [¹¹C]VitC +/- activation with 2 μ M phorbol 12-myristate 13-acetate and +/- 20 μ g/mL cytochalasin B blocking.^{24,15c} For both HL60 cells and neutrophils a significant increase in cell-associated activity, approximately 2-fold, was noted with activation (**p = 0.0025, ***p = 0.00041) (Figure 4b,c). For HL60 cells administration of cytochalasin B decreased the uptake of [¹¹C]VitC to approximately the amount observed for (-) PMA. For neutrophils partial blocking was observed. This partial blocking effect in activated neutrophils could be explained by nitric oxide mediated expression of SVCT, as previously described.²⁶ As expected % cell associated activity of [¹¹C]DHA with co-administration of cytochalasin B did not differ significantly between +/- PMA in human neutrophils due to blocking of GLUT transport (Figure S7). These results provide the basis for detection of endogenously produced ROS using [¹¹C]VitC PET.

In conclusion, we have developed a new PET radiotracer [¹¹C]VitC, which exhibits ROSdependent cellular accumulation. [¹¹C]VitC and its redox partner [¹¹C]DHA behaved as anticipated *in vitro* and *in vivo*, consistent with their markedly different transport mechanisms. [¹¹C]VitC is capable of detecting endogenously produced ROS in activated neutrophil-lineage cells, suggesting potential clinical utility in studying inflammation and/or monitoring immunotherapy. We hypothesize that the ascorbate recycling mechanism may be used to image a myriad of ROS-driven disease states. Furthermore, as high-dose vitamin C has been studied as an anticancer therapy for decades,²⁷ low toxicity in patients has already been well-established.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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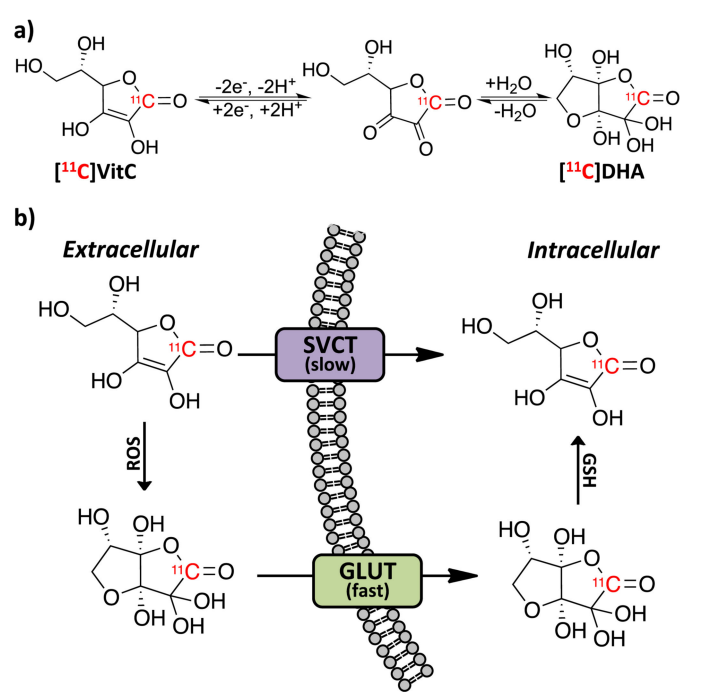


Figure 1.

(a) Oxidation of and hydration of VitC forming bicyclic DHA hydrate. (b) Transport mechanisms of VitC and DHA.

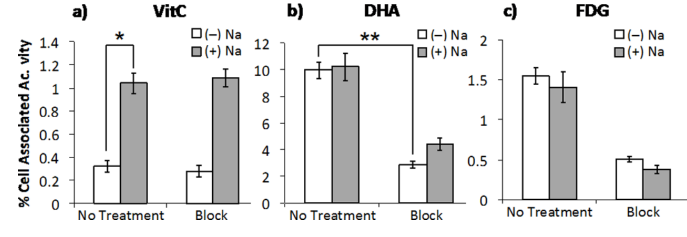


Figure 2.

Uptake of (a) [¹¹C]VitC (*p = 0.0002), (b) [¹¹C]DHA (**p < 0.0001) and (c) [¹⁸F]FDG (+)/(-) availability of Na⁺ in media for co-transport via SVCT, (+)/(-) 10 μ g/mL cytochalasin B blockade of GLUT in U87 human glioblastoma cancer cells.

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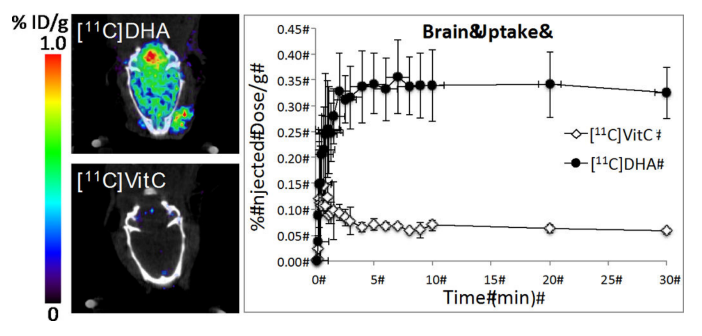
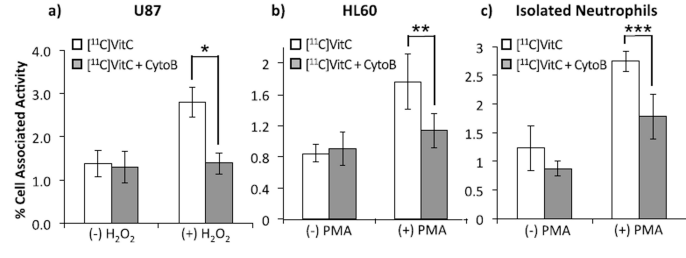


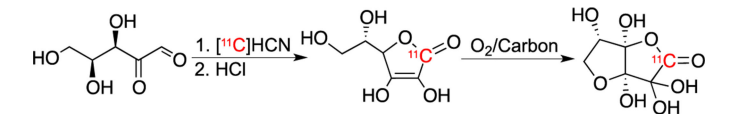
Figure 3.

Representative *in vivo* microPET images of $[^{11}C]$ VitC and $[^{11}C]$ DHA in a normal rat brain (t = 0 - 30 min) and brain ROI data (n = 3) for dynamic scans.





Uptake of [¹¹C]VitC in (a) in U87 glioma cells (+)/(–) 100 μ m H₂O₂, b) HL60 human promeylocyctic leukemia cells (+)/(–) 2 μ M PMA activation and (c) freshly isolated human neutrophils (+)/(–) 2 μ M PMA and (+)/(–) 20 μ g/mL cytochalasin B blockade.



Scheme 1. Radiochemical syntheses of [¹¹C]VitC and [¹¹C]DHA.

Table 1

Summary of radiochemical yields and specific activities for [¹¹C]VitCradiosyntheses with varying amounts of carrier added.

KCN carrier added	% Radiochemical Yield	Specific Activity (mCi/µmol)	number of trials
1 mg/mL	45.4 ± 9.8	4.0 ± 1.2	n = 6
0.75 mg/mL	46.1 ± 9.31	6.4 ± 0.66	n = 3
0.5 mg/mL	29.7 ± 8.8	9.0 ± 3.7	n = 5
0 mg/mL	14.3 ± 10.4	267 ± 148	n = 5